

SEARCH REQUEST FORM

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Requester's Full Name: HELEA Art Unit: 1713 Phone Nu Mail Box and Bldg/Room Location:	1 RETUIO I Imber 30 2 - 1108 REIN - 18 A 2 9 Result	Examiner #: 70058 Date: 62866 Serial Number: 10/081, 628 Examiner #: 70058 Date: 6286 Examiner #: 70088 Date: 6286 Examine					
If more than one search is submit	tted, please prioritize	searches in order of need.					
Please provide a detailed statement of the se Include the elected species or structures, ke utility of the invention. Define any terms the known. Please attach a copy of the cover sh	earch topic, and describe as ywords, synonyms, acronymat may have a special meaneet, pertinent claims, and a	************ specifically as possible the subject matter to be searched. ms, and registry numbers, and combine with the concept or ning. Give examples or relevant citations, authors, etc. if bstract.					
Title of Invention:	ATTACHED						
Inventors (please provide full names):	1						
Earliest Priority Filing Date:	0/22/2001						
For Sequence Searches Only Please includ	e all pertinent information (pe	arent, child, divisional, or issued patent numbers) along with the					
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PTO-1590 (8-01)							



STIC Search Report

STIC Database Tracking Number: 125850

TO: Helen Pezzuto Location: REM 10A29

Art Unit: 1713
July 1, 2004

Case Serial Number: 10/081628

From: Kathleen Fuller Location: EIC 1700 REMSEN 4B28

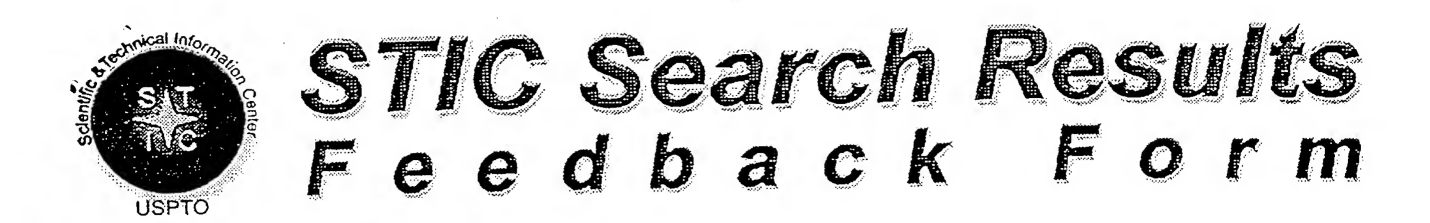
Phone: 571/272-2505

Kathleen.Fuller@uspto.gov

Search Notes

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Questions about the scope or the results of the search? Contact the EIC searcher or contact:

Kathleen Fuller, EIC 1700 Team Leader 571/272-2505 REMSEN 4B28

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 ☐ Foreign Patent(s) ☐ Non-Patent Literature (journal articles, conference proceedings, new product announcements etc.)
 Relevant prior art not found: Results verified the lack of relevant prior art (helped determine patentability). Results were not useful in determining patentability or understanding the invention.
Comments:

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1. (Amended) Phosphorus-containing polymer, suitable for coating dielectric surfaces, of the general formula I or II,

 $P(A)_{m}(F)_{n1}(U)_{o1}$

(I)

 $P(A)_m(UF_{n2})_{o2}$

(II)

in which

hydrophilic polymer

stands for a linear or branched, uncrosslinked or crosslinked, homo- or heteropolymeric polymer component,

(A)

m

stands for identical or different phosphorus-containing groups bonded to P,

A = Ssyp

stands for a number from 3 to 1000,

CF

nl stands for a number from 1 to 1000,

n2 stands for a number from 1 to 100,

stands for identical or different, linear or branched, uncrosslinked or crosslinked oligomeric or polymeric segments, made up of identical or different monomers, which are bonded to P,

- ol stands for a number from 0 to 1000,
- o2 stands for a number from 1 to 1000.

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- 2. (Amended) Polymer according to Claim 1, wherein said polymer contains phosphorus-containing groups A in an amount of from 0.001 to 10 mEq.
- 3. (Amended) Polymer according to Claim 1, wherein said polymer contains functional groups F in an amount of from 0.001 to 20 mEq.
- 4. (Amended) Polymer according to Claim 1, wherein said polymer contains segments U in an amount of from 0.001 to 20 mEq.
- 5. (Amended) Polymer according to Claim 1, wherein the polymer has an average molar mass of from 1000 to 10,000,000 g/mol.
- 6. (Amended) Polymer according to Claim 1, wherein the polymer component P is a statistical copolymer or block copolymer.
- 7. (Amended) Polymer according to Claim 1, wherein the polymer component P is hydrophilic.
- 8. (Amended) Polymer according to Claim 1, wherein said polymer contains phosphorus-containing groups A in the form of a spacer carrying from one to six identical or different phosphorus-containing radicals.

Cont

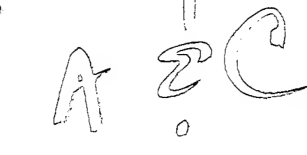
- 9. (Amended) Polymer according to Claim 1, wherein said polymer contains functional groups F that can form covalent bonds, coordination bonds or take part in biochemical recognition reactions.
- 10. (Amended) Polymer according to Claim 1, wherein said polymer contains functional groups F) with crosslinkers.
- 11. (Amended) Polymer according to Claim 1, wherein the segments U have a molar mass, or average molar mass, of from 100 to 10,000.
- 12. (Amended) Polymer according to Claim 1, wherein the groups or segments U are hydrophilic.
- 13. (Amended) Process for preparing a polymer according to Claim 1, comprising the step of copolymerizing
- (A) a monomer containing a phosphorus-containing group A, or a plurality of identical or different monomers containing identical or different phosphorus-containing groups A with
- (B) a monomer containing a functional group F, or a plurality of identical or different monomers containing identical or different functional groups F, and
- (C) optionally, a monomer containing a segment U, or a plurality of identical or different monomers containing identical or different segments U,

to form a polymer of the formula I,

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(B') a monomer containing a unit $(UF_{n2})_{o2}$ according to formula II, or a plurality of identical or different monomers containing identical or different units of the formula $(UF_{n2})_{o2}$ according to formula II,

to form a polymer of the formula II.

- 14. (Amended) Process for preparing a polymer according to Claim 1, comprising the following steps:
- (i) preparing a polymer, which forms the polymer component P and carries identical or different functional groups that are suitable as functional groups F,
- (ii) reacting some of the functional groups to form identical or different phosphorus-containing groups A, and
- (iii) optionally, reacting some of the functional groups to form identical or different segments U, wherein step (iii) can be carried out after, before or together with step (ii), and wherein not all the functional groups are converted in steps (ii) and (iii), and the functional groups that are not converted in steps (ii) and (iii) form the functional groups F of the polymer.
- 15. (Amended) Process according to Claim 14, wherein some or all of the functional groups that have not been converted in steps (ii) and (iii) are reacted with one or more identical or different crosslinkers to form functional groups F.

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16. (Amended) A method of using a polymer according to Claim 1, comprising the step of applying the polymer to a dielectric material so as to form a coating on the dielectric material.

- 17. (Amended) The method of claim 16, wherein the dielectric material is a dielectric waveguide or a portion of a dielectric waveguide.
- 18. (Amended) An optical signal transducer having a coated dielectric waveguide, wherein the coating on the dielectric waveguide consists of a polymer according to claim 1.
- 19. (Amended) A method of using the optical signal transducer of claim 18, wherein said optical signal transducer is exposed to a fluid containing at least one chemical and/or biochemical recognition element which is then immobilized on the coating on the dielectric waveguide.

Please add new claims 20 - 33 as follows.

20. (New) Polymer according to Claim 1, wherein said polymer contains phosphorus-containing groups A in an amount of from 0.01 to 5 mEq.

- 21. (New) Polymer according to Claim 1, wherein said polymer contains phosphorus-containing groups A in an amount of from 0.1 to 3 mEq.
- 22. (New) Polymer according to Claim 1, wherein said polymer contains functional groups F in an amount of from 0.01 to 10 mEq.

23. (New) Polymer according to Claim 1, wherein said polymer contains functional groups F in an amount of from 0.5 to 10 mEq.

- 24. (New) Polymer according to Claim 1, wherein said polymer contains segments U in an amount of from 0.01 to 10 mEq.
- 25. (New) Polymer according to Claim 1, wherein said polymer contains segments U in an amount of from 0.5 to 10 mEq.
- 26. (New) Polymer according to Claim 1, wherein the polymer has an average molar mass of from 2100 to 1,000,000 g/mol.
- 27. (New) Polymer according to Claim 1, wherein the polymer has an average molar mass of from 5000 to 500,000 g/mol.
- 28. (New) Polymer according to Claim 1, wherein the polymer has an average molar mass of from 5000 to 300,000 g/mol.
- 29. (New) Polymer according to Claim 1, wherein the polymer has an average molar mass of from 10,000 to 150,000 g/mol.
- 30. (New) Process for preparing a polymer according to Claim 1, comprising the following steps:
- (i) preparing a polymer, which forms the polymer component P and carries identical or different functional groups that are suitable as functional groups F, said functional groups F being selected from the group consisting of hydroxyl groups, carboxyl groups, derivatives of carboxyl groups and amine groups,
- (ii) reacting some of the functional groups to form identical or different phosphorus-containing groups A, and

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(iii) optionally, reacting some of the functional groups to form identical or different segments U, wherein step (iii) can be carried out after, before or together with step (ii), and wherein not all the functional groups are converted in steps (ii) and (iii), and the functional groups that are not converted in steps (ii) and (iii) form the functional groups F of the polymer.

- 31. (New) The method of claim 16, wherein the dielectric material comprises at least one material selected from the group consisting of TiO₂, Ta₂O₅, ZrO₂, HfO₂ and Al₂O₃.
- 32. (New) The method of claim 16, wherein the dielectric material comprises at least one material selected from the group consisting of TiO₂ and Ta₂O₅.
- 33. (New) An optical signal transducer having a coated dielectric waveguide, wherein the coating on the dielectric waveguide consists of a polymer according to claim 1 and the dielectric waveguide comprises at least one material selected from the group consisting of TiO₂, Ta₂O₅, ZrO₂, HfO₂ and

s stands for the number 1 (i.e. $A = SY_p$)

and in which the group or groups Y is are selected from the following phosphorus-containing radicals:

 $-O(R'O)PO_2H, -P(R'O)O_2H, -N(CH_2-P(R'O)O_2H)_2, -N(R')-CH_2-P(R'O)O_2H, -CH(P(R'O)O_2H)N(CH_2-P(R'O)O_2H)_2, -CH(CH_2-P(R'O)O_2H)_2, -CR'(CH_2-P(R'O)O_2H)_2, -C(CH_2-P(R'O)O_2H)_3,$

where R' stands for -H, -CH₃ or -C₂H₅.

The polymer preferably contains one or more of the following groups Y:

-O(R'O)PO₂H, -P(R'O)O₂H, -N(CH₂-P(R'O)O₂H)₂, in particular -N(CH₂-P(R'O)O₂H)₂,

where R' preferably stands for -H.

The spacer S is directly coupled to a C atom of the polymer and carries p identical or different phosphorus-containing radicals Y. According to the invention, the following spacers are preferred (group(s) Y are also indicated):

-(CH₂)_q-(O-CH₂-CH₂)_r-Y, -(CH₂)_q-(O-CH₂-CH₂-CH₂)_r-Y, -(CH₂)_q-(O-CH₂-CH₂)_r-C₆H₄Y, -(CH₂)_q-(O-CH₂-CH₂)_r-C₆H₃Y₂, -(CH₂)_q-(O-CH₂-CH₂)_r-C₆H₂Y₃,

where q stands for numbers from 0 to 20 and r stands for numbers from 20 0 to 100.

In a particular embodiment, the polymer according to the invention contains phosphorus-containing groups A in the form of a spacer S carrying from one to six identical or different phosphorus-containing radicals.

The following groups A, which are coupled directly to a C atom of the polymer, are preferred:

-PO₃H₂, -NH-CH₂-CH₂-PO₃H₂, -CH₂-N(CH₂-PO₃H₂)₂, -N(CH₂-PO₃H₂)₂, -(CH₂)₄N(CH₂-PO₃H₂)₂, -OPO₃H₂.

Functional groups F for immobilizing recognition elements

F stands for functional groups which are bonded directly to a carbon atom of the polymer, and via which recognition elements can be immobilized directly or with the aid of a crosslinker covalently, coordinatively or via another chemical bond onto the polymer, and therefore onto the surface of the bio- or chemosensor. The direct coupling of the recognition elements can be carried out

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Examples

Example 1: Polymer made from phosphonate-functional copolymers.

A mixture of 50 g of N-methyl-2-pyrrolidone (NMP), 5 g of vinylphosphonic acid, 10 g of triethylamine, 15 g of methacryloxyethyl acetoacetate, 30 g of polyethylene glycol methyl ether acrylate (molar mass 750 g/mol), 0.5 g of azobisisobutyronitrile and 1.5 g of dodecyl mercaptan was heated for 6 h to 65°C. After cooling, the solution was adjusted in ethanol to a concentration of 0.1 mg of polymer per ml of solution, and the waveguide surfaces were incubated in this solution for 18 h. The waveguides were then washed with ethanol and 10 mM (M = mol/l) NaOH. A solution of 2 mg/ml of anti-myoglobin monoclonal mouse antibodies in 10 mM sodium acetate buffer, adjusted to pH = 5, was prepared and the waveguide surfaces were incubated in it for 2 h. A 1.5 ng/mm² surface concentration of antibodies was obtained.

Example 2: Polymer made from phosphate esters of polyvinyl alcohol.

A mixture of 50 g of a 10% strength solution of polyvinyl alcohol (polyvinyl acetate with an 88% degree of saponification and a Höppler viscosity of 18 for the 4% strength solution in water) in DMSO and 0.1% of polyphosphoric acid was heated for 15 min to 100°C. After cooling, 10 g of succinic anhydride were added to the solution and stirred at 21°C for 3 h. In the next step, the solution was adjusted in ethanol to a concentration of 1 mg of polymer per ml of solution, and the waveguide surfaces were incubated in this solution for 18 h. The waveguides were then washed with ethanol and 10 mM NaOH. The surface was incubated for 10 min in a solution of 1 M N-hydroxysuccinimide and 1 M N-dimethylaminopropyl-N'-ethyl-carbodiimide hydrochloride in ultrapure water, and then washed with ultrapure water. A solution of 2 mg/ml of anti-myoglobin monoclonal mouse antibodies in 10 mM sodium acetate buffer, adjusted to pH = 5, was prepared and the waveguide surfaces were incubated in it for 2 h. A 2.5 ng/mm² surface concentration of antibodies was obtained.

Example 3: Polymer made from imidised MSA copolymers.

15.6 g of polymaleic anhydride-alt-methyl vinyl ether MW (average molar mass) = 216,000 g/mol) were added portionwise to a mixture of 9.5 g of 2-

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(2-aminoethoxy)-ethanol, 1.11 g of aminomethanephosphonic acid, 1 g of triethylamine and 100 ml of water at 70°C. After cooling, the solution was adjusted in ethanol to a concentration of 10 mg of polymer per ml of solution, and the waveguide surfaces were incubated in this solution for 18 h. The waveguides were then washed with ethanol and 10 mM NaOH. The surfaces were incubated in a 10 mg/ml solution of ethylene glycol bissuccinimidyl succinate in DMSO for 30 min and then washed with DMSO and ultrapure water. A solution of 2 mg/ml of anti-human chorionic gonadotropin monoclonal mouse antibodies in 10 mM sodium acetate buffer, adjusted to pH = 5, was prepared and the waveguide surfaces were incubated in it for 2 h. A 2.0 ng/mm² surface concentration of antibodies was obtained.

Example 4: Polymer made from phosphonate-functional copolymers grafted with polyglycidol.

Preparation of the grafting basis (polyglycidol modified with fatty acid):

A mixture of 28 g of soybean oil fatty acid and 74 g of epoxypropanol (glycidol) was heated for 1 h to 140°C and then a mixture of 0.4 g of phosphoric acid and 333.5 g of epoxypropanol was added in portions over 6 h. The mixture was then stirred for a further 16 h at 140°C.

A mixture of 20 g of the previously prepared polyglycidol modified with fatty acid, 20 g of methacryloyloxyethyl acetoacetate, 2 g of vinylphosphonic acid, 2 g of triethylamine, 42 g of NMP and 0.4 g of azobisisobutyronitrile was heated for 16 h to 65°C and for 1 h to 100°C. After cooling, the solution was adjusted in ethanol to a concentration of 3 mg of polymer per ml of solution, and the waveguide surfaces were incubated in this solution for 10 h. The waveguides were then washed with ethanol and 10 mM NaOH. A solution of 2 mg/ml of antimyoglobin monoclonal mouse antibodies in 10 mM sodium acetate buffer, adjusted to pH = 5, was prepared and the waveguide surfaces were incubated in it for 2 h. A 3.5 ng/mm² surface concentration of antibodies was obtained.

Example 5: Polyglycidol, derivatized with maleic acid anhydride and imino-bis-methylene phosphonic acid.

Preparation of the thiol derivatized imido-di-methylene phosphonic acid reagent:

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A mixture of 100 g of mercapto ethylamine hydrochloride, 150 g phosphonic acid and 170 g of water was heated to 100°C and over 1 h 287 g of formaldehyde (37% strength) were added dropwise. The mixture was stirred for a further hour and then the solvent was removed under vacuum.

Preparation of the polyglycidol:

1.88 g of hexadecyl amine were melted in a 250 ml glas reactor heated to 100°C and reacted with 1.2 g glycidol. Then 0.9 ml of potassium ethoxide solution (25% strength in methanol) was added and excessive methanol removed under vacuum. At 140°C the residue was dissolved in 15 ml of dry diglyme. At a speed of 25 ml per hour 260 g of glycidol in 350 ml of dry THF were added in portions. Upon completion of the addition the reaction mixture was dissolved in 1200 ml of methanol and neutralized by filtration over an acidic ion exchanger (Amberlite® IR-120). The filtrate was precipitated in 121 of acetone and the yielded polymer was dried at 80°C for 12 h under vacuum. 254 g of a colorless, highly viscous liquid with a molar mass of 30,000 g/mol and a polydispersity of 1.23 were received. All molecules comprise the initiator as Kerneinheit and 27% of branched building units.

Subsequently a mixture of 1 g of the previously prepared polyglycidol and 5 g of DMSO was heated to 50°C. Then 0.2 g of maleic acid anhydride was added. After 15 min it was heated to 80°C and 0.2 g of thiol derivatized imido-bismethylene phosphonic acid reagent and 0.3 g triethyl amine were added. After 15 min 0.05 g of azoisobutyro nitrile was added and it was stirred for a further 4 h at 80°C and then for a further hour at 100°C.

After cooling, the solution was adjusted in ethanol to a concentration of 2 mg of polymer per ml of solution, and the waveguide surfaces were incubated in this solution for 16 h. The waveguides were then washed with ethanol and water. The surface was incubated in a solution of 1 M of N-hydroxy succinimide and 1 M of N-dimethyl aminopropyl N'-ethyl carbodiimide hydrochloride in ultrapure water for 10 min and then washed with ultrapure water. A solution of 2 mg/ml of anti-myoglobin monoclonal mouse antibodies in 10 mM sodium acetate buffer, adjusted to pH = 5, was prepared and the waveguide surfaces were incubated in it for 2 h. A 2.8 ng/mm² surface concentration of antibodies was obtained.

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Example 6: Polymer made from dextran modified with acetoacetoxy and phosphate ester.

A mixture of 10 g of dextran (MW = 40,000 g/mol), 7 g of tert-butyl acetoacetate, 100 g of DMSO and 0.5 g of polyphosphoric acid was heated for 4 h to 80°C. After cooling, the solution was adjusted in ethanol to a concentration of 1 mg of polymer per ml of solution, and the waveguide surfaces were incubated in this solution for 8 h. The waveguides were then washed with ethanol and 10 mM NaOH. A solution of 2 mg/ml of streptavidin in 10 mM sodium acetate buffer, adjusted to pH = 5, was prepared and the waveguide surfaces were incubated in it for 2 h. A 4.5 ng/mm² surface concentration of streptavidin was obtained.

Example 7: Polymer made from phosphonate-functional polylysine.

500 mg of poly-L-lysin hydrobromide (MW = 150,000 to 300,000 g/mol), 170 mg of phosphoric acid and 4 ml of water was heated to 100°C, and then 324 mg of formaldehyde (37% strength) were added. The mixture was stirred for 1 h at 100°C. After cooling, the solution was adjusted in ethanol to a concentration of 1 mg of polymer per ml of solution, and the waveguide surfaces were incubated in this solution for 2 h. The waveguides were then washed with ethanol and 10 mM NaOH. The surfaces were incubated with a solution of 10 mg/ml of carboxymethyldextran (MW = 15,000 g/mol), 0.1 M of N-hydroxysuccinimide and 0.1 M of N-dimethylaminopropyl-N-ethylcarbodiimide hydrochloride in ultrapure water for 20 min. The surfaces were then washed briefly with ultrapure water and immediately incubated with 0.1 mg/ml of an amine-functionalized DNA (20 nucleotides) in 10 mM sodium acetate buffer (pH = 5). A 0.5 ng/mm² surface concentration of DNA was obtained.

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L5	117070	SEA FILE=REGISTRY SSS FUL L1 NOT L3
L6	9950	SEA FILE=REGISTRY ABB=ON L5 AND PMS/CI
L7	8527	SEA FILE=HCAPLUS ABB=ON L6
L8	4271	SEA FILE=HCAPLUS ABB=ON L5/DP
L9	378	SEA FILE=HCAPLUS ABB=ON L8(L)POLYMER?
L10	3	SEA FILE=HCAPLUS ABB=ON L9 AND ?SENSOR?
L11	1	SEA FILE=HCAPLUS ABB=ON L9 AND (TRANDUC? OR WAVEGUIDE? OR
		WAVE(W)GUIDE?)
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		SEA FILE=HCAPLUS ABB=ON L8(L)?POLYMER?
	272	SEA FILE=HCAPLUS ABB=ON L8 AND POLYMER?/SC,SX
L15	10	SEA FILE=HCAPLUS ABB=ON (L13 OR L14) AND (TRANSDUC? OR
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L16	10	SEA FILE=HCAPLUS ABB=ON L15 OR L10 OR L11 OR L12
L17	200	SEA FILE=HCAPLUS ABB=ON L7 AND (TRANSDUC? OR DIELECTRIC? OR
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L18	2378	SEA FILE=HCAPLUS ABB=ON L7(L)(PREP OR IMF OR SPN)/RL
L19		SEA FILE=HCAPLUS ABB=ON L17 AND L18
L20	9	SEA FILE=HCAPLUS ABB=ON L19 AND BIOCHEM?/SC,SX
L21	42	SEA FILE=HCAPLUS ABB=ON L7(L)(TRANSDUC? OR DIELECTRIC? OR
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Conjugation of enzymes on polymer nanoparticles covered with ${
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phosphorylcholine groups Konno, T.; Watanabe, J.; Ishihara, K. ΑU Department of Materials Engineering, School of Engineering, The University CS of Tokyo, Bunkyo, Tokyo, 113-8656, Japan Biomacromolecules (2004), 5(2), 342-347 SO CODEN: BOMAF6; ISSN: 1525-7797 American Chemical Society PB DT Journal English LACC 9-16 (Biochemical Methods) Section cross-reference(s): 7 We investigated the bioconjugation of enzymes on polymer nanoparticles ABcovered with bioinert phosphorylcholine groups. A water-soluble amphiphilic phospholipid polymer (PMBN) was specially designed for preparation of nanoparticles and conjugation with enzymes on them. The PMBN was prepared by random copolymn. of 2-methacryloyloxyethyl phosphorylcholine (MPC), Bu methacrylate, and p-nitrophenylester bearing methacrylate. The PMBN was used as an emulsifier and a surface modifier to prepare the poly(L-lactic acid) nanoparticles by a solvent evaporation technique in aqueous medium. nanoparticles covered with phosphorylcholine groups were stably dispersed in an aqueous solution and a phosphate buffered saline. The diameter and surface ζ -potential of the nanoparticles were ca. 200 Nm and -6 mV, resp. The p-nitrophenyl ester groups, which are active ester units for the amino groups of the protein, were located at the surface of the nanoparticles. Both acetylcholine esterase and choline oxidase were co-immobilized (dual-mode conjugation) by the reaction between the p-nitrophenyl ester group and the amino group of these enzymes. The enzymic reactions on the nanoparticles were followed using a microdialysis biosensor system with a microtype hydrogen peroxide electrode in the probe. The nanoparticles conjugated with these enzymes could detect the acetylcholine chloride as hydrogen peroxide, which is a product of the enzymic reactions on the surface of the nanoparticles in the probe. Namely, continuous enzyme reactions could be occurring on the surface of the nanoparticles. It is concluded that the nanoparticles are a promising tool for a highly sensitive and microdiagnostic system. conjugation enzyme polymer nanoparticle covered phosphorylcholine group STAtomic force microscopy ITConjugation (molecular association) Fluorometry Nanoparticles Zeta potential (conjugation of enzymes on polymer nanoparticles covered with phosphorylcholine groups) 60-31-1, Acetylcholine chloride 67-48-1, Choline chloride 7722-84-1, ITHydrogen peroxide, analysis RL: ANT (Analyte); ANST (Analytical study) (conjugation of enzymes on polymer nanoparticles covered with phosphorylcholine groups) 9000-81-1, E.C.3.1.1.7 9028-67-5, E.C.1.1.3.17 IT RL: ARU (Analytical role, unclassified); PEP (Physical, engineering or chemical process); PYP (Physical process); ANST (Analytical study); PROC (Process) (conjugation of enzymes on polymer nanoparticles covered with phosphorylcholine groups) IT666711-04-2P RL: ARU (Analytical role, unclassified); PEP (Physical, engineering or

The

chemical process); PYP (Physical process); SPN (Synthetic preparation); ANST (Analytical study); PREP (Preparation); PROC (Process)

(conjugation of enzymes on polymer nanoparticles covered with phosphorylcholine groups)

IT 666711-01-9

RL: RCT (Reactant); RACT (Reactant or reagent)

(conjugation of enzymes on polymer nanoparticles covered with phosphorylcholine groups)

RE.CNT 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD RE

- (1) Davies, M; Adv Drug Delivery Rev 2000, V45, P169 HCAPLUS
- (2) Fisher, A; Alzheimer's and Parkinson's Diseases: Strategies for Research and Development 1986
- (3) Ishihara, K; Colloid Surf B: Biointerfaces 2000, V18, P325 HCAPLUS
- (4) Ishihara, K; J Biomed Mater Res 1990, V24, P1069 HCAPLUS
- (5) Ishihara, K; J Biomed Mater Res 1992, V26, P1543 HCAPLUS
- (6) Ishihara, K; Polym Adv Technol 2000, V11, P626 HCAPLUS
- (7) Ishihara, K; Polym J 1990, V22, P355 HCAPLUS
- (8) Ishihara, K; Polym J 1999, V31, P1231 HCAPLUS
- (9) Iwasaki, Y; J Biomed Mater Res 2001, V57, P74
- (10) Konno, T; Biomaterials 2001, V22, P1883 HCAPLUS
- (11) Larsson, N; Electrochim Acta 1998, V43, P3541 HCAPLUS
- (12) Solovskij, M; Eur Polym J 2000, V36, P1127 HCAPLUS
- (13) Torchilin, V; Biochim Biophy Acta 2001, V1511, P397 HCAPLUS
- (14) Torto, N; Anal Chim Acta 1999, V379, P281 HCAPLUS
- (15) Trubetskoy, V; Adv Drug Delivery Rev 1999, V37, P81 HCAPLUS

IT 666711-04-2P

RL: ARU (Analytical role, unclassified); PEP (Physical, engineering or chemical process); PYP (Physical process); SPN (Synthetic preparation); ANST (Analytical study); PREP (Preparation); PROC (Process)

(conjugation of enzymes on polymer nanoparticles covered with phosphorylcholine groups)

RN 666711-04-2 HCAPLUS

CN 3,5,8-Trioxa-4-phosphaundec-10-en-1-aminium, 4-hydroxy-N,N,N,10-tetramethyl-9-oxo-, inner salt, 4-oxide, polymer with butyl 2-methyl-2-propenoate and α -(2-methyl-1-oxo-2-propenyl)- ω -[[(4-nitrophenoxy)carbonyl]oxy]poly(oxy-1,2-ethanediyl), graft (9CI) (CA INDEX NAME)

CM 1

CRN 666711-01-9

CMF (C2 H4 O)n C11 H9 N O6

CCI PMS

CM 2

CRN 67881-98-5

CMF C11 H22 N O6 P

CM 3

CRN 97-88-1 CMF C8 H14 O2

L23 ANSWER 2 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 2004:139242 HCAPLUS

DN 140:207492

ED Entered STN: 20 Feb 2004

TI (Copy)Oil-based platemaking electrostatic ink-jet printing ink containing star block copolymer dispersion stabilizing resin for improved ink ejection stability

IN Kato, Eiichi

PA Fuji Photo Film Co., Ltd., Japan

SO Jpn. Kokai Tokkyo Koho, 57 pp. CODEN: JKXXAF

DT Patent

LA Japanese

IC ICM C09D011-00

ICS B41J002-01; B41M005-00; B41C001-10

CC 74-6 (Radiation Chemistry, Photochemistry, and Photographic and Other Reprographic Processes)

Section cross-reference(s): 35, 38

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

PI JP 2004051787 A2 20040219
PRAI JP 2002-211352 20020719

AB The oil-based platemaking electrostatic ink-jet printing ink comprises a chargeable resin particle dispersed in a nonaq. support solution having the elec. resistivity $\geq 109~\Omega \cdot \text{cm}$ and the **dielec**.

constant ≤3.5. The resin particle is obtained from (A) ≥1 monofunctional monomer which is soluble in an nonag. solvent but

monofunctional monomer which is soluble in an nonaq. solvent but becomes soluble

upon polymn, (B) ≥1 monofunctional monomer which is copolymerizable with said monomer (A) and has an acid group, ≥1 monofunctional macromer having a double bond at the end group copolymerizable with said monomer (A), having F- and/or Si-containing repeating unit, and having the weight

average mol. weight $\leq 20,000$, and a star-type copolymer dispersion stabilizing resin having ≥ 3 sp. block copolymer chains bonded to an

organic residue. oil platemaking electrostatic ink jet printing star block copolymer; STlithog printing plate ink dispersion stabilizing resin Acrylic polymers, uses ΙT RL: TEM (Technical or engineered material use); USES (Uses) (F-containing; oil-based platemaking electrostatic ink-jet printing ink containing star block copolymer dispersion stabilizing resin for improved ink ejection stability) ITInks (jet-printing; oil-based platemaking electrostatic ink-jet printing ink containing star block copolymer dispersion stabilizing resin for improved ink ejection stability) Lithographic plates IT(oil-based platemaking electrostatic ink-jet printing ink containing star block copolymer dispersion stabilizing resin for improved ink ejection stability) 150551-83-0 150551-89-6 150551-92-1 150551-93-2 150551-97-6 IT154340-06-4 155161-59-4 155293-25-7 159967-38-1 159967-39-2 159967-41-6 159967-42-7 159967-43-8 216877-91-7 RL: RCT (Reactant); RACT (Reactant or reagent) (core; preparation of star block copolymer for oil-based platemaking electrostatic ink-jet printing ink) 150469-59-3P 159967-35-8P 159967-45-0P 159967-46-1P 159967-47-2P IT159967-48-3P 159967-49-4P 159967-50-7P 159967-51-8P 159967-53-0P 159967-54-1P 159967-55-2P 159967-56-3P 216878-23-8P 524745-45-7P RL: PRP (Properties); SPN (Synthetic preparation); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses) (oil-based platemaking electrostatic ink-jet printing ink containing star block copolymer dispersion stabilizing resin for improved ink ejection stability) 96-33-3DP, Methyl acrylate, F-containing acrylic polymer 140-88-5DP, Ethyl ITacrylate, F-containing acrylic polymer 2495-37-6DP, Benzyl methacrylate, F-containing acrylic polymer 524745-47-9DP, F-containing acrylic polymer 524745-88-8DP, F-containing acrylic polymer 524745-90-2DP, F-containing acrylic polymer 524745-92-4DP, F-containing acrylic polymer 524745-94-6DP, F-containing acrylic polymer 524745-96-8DP, F-containing acrylic polymer 524745-98-0DP, F-containing acrylic polymer 661482-71-9DP, F-containing acrylic polymer RL: SPN (Synthetic preparation); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses) (oil-based platemaking electrostatic ink-jet printing ink containing star block copolymer dispersion stabilizing resin for improved ink ejection stability) 159967-36-9P, Methyl acrylatemethyl methacrylate-octadecyl methacrylate ${ t IT}$ block copolymer RL: PRP (Properties); SPN (Synthetic preparation); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses) (star; oil-based platemaking electrostatic ink-jet printing ink containing star block copolymer dispersion stabilizing resin for improved ink ejection stability) 524745-90-2DP, F-containing acrylic polymer IT524745-94-6DP, F-containing acrylic polymer 524745-96-8DP, F-containing acrylic polymer 524745-98-0DP, F-containing acrylic polymer 661482-71-9DP, F-containing acrylic polymer RL: SPN (Synthetic preparation); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses) (oil-based platemaking electrostatic ink-jet printing ink containing star

block **copolymer** dispersion stabilizing resin for improved ink ejection stability)

RN 524745-90-2 HCAPLUS

CN Methanol, [[(4-ethenylphenyl)methyl]octylamino]-, dihydrogen phosphate (ester) (9CI) (CA INDEX NAME)

$$H_2O_3PO-CH_2$$
 $Me-(CH_2)_7-N-CH_2$

RN 524745-94-6 HCAPLUS

CN 2-Propenoic acid, 2-methyl-, 2-[[(3-phosphonophenyl)methyl]amino]ethyl ester (9CI) (CA INDEX NAME)

$$H_{2}O_{3}P$$
 $CH_{2}-NH-CH_{2}-CH_{2}-O-C-C-Me$

RN 524745-96-8 HCAPLUS

CN 2-Propenamide, N-[3-[[(phosphonooxy)methyl]propylamino]propyl]- (9CI) (CA INDEX NAME)

$$CH_2-OPO_3H_2$$
 O | | n-Pr-N-(CH₂)3-NH-C-CH= CH₂

RN 524745-98-0 HCAPLUS

CN Butanoic acid, 4-phosphono-, 1-[2-amino-3-[(2-methyl-1-oxo-2-propenyl)oxy]propyl] ester (9CI) (CA INDEX NAME)

RN 661482-71-9 HCAPLUS

CN 2-Propenoic acid, 2-methyl-, 2-[[2-(diethylamino)ethyl][(phosphonooxy)methyl]amino]ethyl ester (9CI) (CA INDEX NAME)

L23 ANSWER 3 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

KATHLEEN FULLER EIC 1700 REMSEN 4B28 571/272-2505

```
2004:134082 HCAPLUS
AN
     140:190011
DN
    Entered STN: 19 Feb 2004
ED
     Oil-based ink for platemaking by electrostatic ink-jet printing method
TI
     Kato, Eiichi
ΙN
     Fuji Photo Film Co., Ltd., Japan
PA
     Jpn. Kokai Tokkyo Koho, 66 pp.
SO
     CODEN: JKXXAF
     Patent
\mathsf{DT}
LA
     Japanese
IC
     ICM C09D011-00
     ICS B41M005-00
    74-6 (Radiation Chemistry, Photochemistry, and Photographic and Other
CC
     Reprographic Processes)
     Section cross-reference(s): 42
FAN. CNT 1
     PATENT NO.
                      KIND DATE
                                           APPLICATION NO. DATE
    JP 2004051763 A2
                                           JP 2002-210572
                            20040219
                                                             20020719
PΙ
PRAI JP 2002-210572
                            20020719
     The ink contains chargeable resin particles dispersed in a water-free
AB
     medium having elec. resistivity \geq 109 \Omega-cm and dielec
     . constant ≤3.5, wherein the resin particles are obtained by polymerizing
     nonaq. solns. containing (A) nonaq. solvent-soluble monofunctional monomers
which
     become insol. after polymerization, (B) NR1R2- and PO3H2- and/or
SO3H-containing
     monofunctional comonomers (R1, R2 = H, C1-22 hydrocarbyl, R1 and R2 may
     form ring), (MM) copolymerizable monofunctional micromonomers having main
     chains of F- and/or Si-containing substituent-containing repeating
unit-containing
     polymers terminated for one side with defined polymerizable double bonds
     and weight-average mol. weight \leq 2 + 104, and (P) nonaq. solvent-soluble
     dispersant polymers containing [Cb1HCb2(V0L)] [V0 = CO2, OCO, (CH2)rCO2,
     (CH2)rOCO, O, C6H4X; X = direct bond, O, OCO, CO2; r = 1-12; L = C8-32
     alkyl or alkenyl, defined siloxane structure- or silyloxy-containing Si1-30
     substituent; b1, b2 = H, halo, cyano, C1-7 hydrocarbyl, CO2H, etc.] in
     which main chains of the polymers are partially crosslinked. The ink
     shows stable delivery and forms clear image with high strength for the
     printing plate with high printability.
    platemaking electrostatic ink jet printing oily ink; polymer particle oil
ST
     based jet printing ink
     Fluoropolymers, preparation
\operatorname{IT}
     Polysiloxanes, preparation
     RL: IMF (Industrial manufacture); TEM (Technical or engineered material
     use); PREP (Preparation); USES (Uses)
        (acrylic, graft, particles; oil-based ink containing polymer particles for
        platemaking by electrostatic ink-jet printing method)
ΙT
     Inks
        (jet-printing; oil-based ink containing polymer particles for platemaking
        by electrostatic ink-jet printing method)
    Lithographic plates
IT
        (oil-based ink containing polymer particles for platemaking by
        electrostatic ink-jet printing method)
IT
     Macromonomers
     RL: IMF (Industrial manufacture); RCT (Reactant); PREP (Preparation); RACT
```

(oil-based ink containing polymer particles for platemaking by

KATHLEEN FULLER EIC 1700 REMSEN 4B28 571/272-2505

electrostatic ink-jet printing method)

(Reactant or reagent)

Dispersing agents

IT

```
(polymeric; oil-based ink containing polymer particles for platemaking by
       electrostatic ink-jet printing method)
\operatorname{IT}
    660426-31-3DP, trimethylsilyl ether
    RL: IMF (Industrial manufacture); TEM (Technical or engineered material
    use); PREP (Preparation); USES (Uses)
        (comprised of actual and assumed monomers, particles; oil-based ink
       containing polymer particles for platemaking by electrostatic
       ink-jet printing method)
    920-46-7DP, Methacrylic chloride, ester with azobiscyanopentanol-modified
IT
    ethylene glycol diacrylate-octadecyl acylate copolymer
    4,4'-Azobis(4-cyanopentanol), reaction products with ethylene glycol
    diacrylate-octadecyl acrylate copolymer, ester with methacrylic chloride
    5926-95-4DP, Glutaconic anhydride, reaction products with
    divinylbenzene-mercaptoethylamine-octadecyl methacrylate telomer
    61255-17-2P, Dodecyl methacrylate-divinylbenzene copolymer 122324-74-7P,
    Divinylbenzene-octadecyl methacrylate copolymer 130805-21-9P,
    Divinylbenzene-tridecyl methacrylate copolymer 139720-57-3P
    139720-64-2DP, Divinylbenzene-2-mercaptoethylamine-octadecyl methacrylate
    telomer, reaction products with glutaconic anhydride 148532-67-6P,
    Dodecyl methacrylate-octyl methacrylate-trivinylbenzene copolymer
    148640-01-1P, Divinylbenzene-octadecyl methacrylate-thioglycolic acid
    telomer ester with 2-hydroxyethyl methacrylate 159446-42-1P
                   214772-24-4P, Divinylbenzene-2-mercaptoethanol-octadecyl
    methacrylate telomer ester with 2-carboxyethyl acrylate 214772-26-6P,
    Divinylbenzene-2-mercaptoethanol-octadecyl methacrylate telomer ester with
    α-chloroacrylic acid
                           214772-29-9P 218459-53-1P, Allyl
    methacrylate-dodecyl methacrylate-thioglycolic acid telomer ester with
    2-hydroxyethyl methacrylate 218459-75-7P 218459-76-8P
                                                                218459-77-9DP.
    Ethylene glycol diacrylate-octadecyl acrylate copolymer, reaction products
    with azobiscyanopentanol, ester with methacrylic chloride
                                                                308283-76-3P,
    Docosyl methacrylate-polyethylene glycol diacrylate copolymer
                  657408-63-4P
                                  657408-64-5P 657408-65-6P
     524745-38-8P
                                                                658039-85-1P
                   660431-29-8P
     660426-29-9P
    RL: IMF (Industrial manufacture); TEM (Technical or engineered material
    use); PREP (Preparation); USES (Uses)
        (dispersant; oil-based ink containing polymer particles for platemaking by
       electrostatic ink-jet printing method)
    312260-55-2P 312260-57-4P 312260-79-0P 312260-82-5P 312260-85-8P
    312260-87-0P 312260-89-2P
                                  312260-91-6P
                                                 312260-93-8P
                                                                312260-96-1P
                                  312261-21-5P 312261-24-8P
                                                                312261-27-1P
    312261-02-2P 312261-17-9P
    312261-30-6P 477719-06-5DP, butanoic acid terminated, ester with
    glycidyl methacrylate 660431-40-3P
    RL: IMF (Industrial manufacture); RCT (Reactant); PREP (Preparation); RACT
     (Reactant or reagent)
        (macromer; oil-based ink containing polymer particles for platemaking by
       electrostatic ink-jet printing method)
    557799-78-7P 557799-79-8P 557799-80-1P 660426-30-2P
\operatorname{IT}
                                                                660426-32-4P
    660426-33-5P 660426-34-6P 660426-37-9P 660426-38-0P
                                                                660426-39-1P
    660426-40-4P 660426-41-5P 660426-42-6P 660426-43-7P
                                                                660426-44-8P
    660426-45-9P 660426-46-0P 660426-47-1P 660426-48-2P
                                                                660426-49-3P
    660426-50-6P 660426-51-7P 660426-52-8P 660426-53-9P
    RL: IMF (Industrial manufacture); TEM (Technical or engineered material
    use); PREP (Preparation); USES (Uses)
        (particles; oil-based ink containing polymer particles for platemaking by
       electrostatic ink-jet printing method)
    660426-31-3DP, trimethylsilyl ether
IT
    RL: IMF (Industrial manufacture); TEM (Technical or engineered material
    use); PREP (Preparation); USES (Uses)
```

(comprised of actual and assumed monomers, particles; oil-based ink containing **polymer** particles for platemaking by electrostatic ink-jet printing method)

RN 660426-31-3 HCAPLUS

CN 2-Propenoic acid, 2-methyl-, 4-[ethyl[(phosphonooxy)methyl]amino]butyl ester, polymer with dimethylsilanediol, methyl 2-methyl-2-propenoate and methyl 2-propenoate, graft (9CI) (CA INDEX NAME)

CM 1

CRN 524745-45-7 CMF C11 H22 N O6 P

CM 2

CRN 1066-42-8 CMF C2 H8 O2 Si

CM 3

CRN 96-33-3 CMF C4 H6 O2

CM 4

CRN 80-62-6 CMF C5 H8 O2

L23 ANSWER 4 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

KATHLEEN FULLER EIC 1700 REMSEN 4B28 571/272-2505

```
AN
     2002:807565 HCAPLUS
DN
     137:331033
ED
     Entered STN: 23 Oct 2002
     Liquid developer containing dispersion resin particles for
TI
     electrophotographic printing platemaking system
     Kato, Eiichi
ΙN
     Fuji Photo Film Co., Ltd., Japan
PA
     Jpn. Kokai Tokkyo Koho, 37 pp.
SO
    CODEN: JKXXAF
\mathsf{DT}
     Patent
LA
    Japanese
IC
     ICM G03G009-13
CC
    74-3 (Radiation Chemistry, Photochemistry, and Photographic and Other
     Reprographic Processes)
     Section cross-reference(s): 35, 38
FAN.CNT 1
     PATENT NO.
                     KIND DATE
                                           APPLICATION NO. DATE
    JP 2002311654 A2
                                           JP 2001-111587 20010410
                           20021023
ΡI
PRAI JP 2001-111587
                            20010410
    The liquid developer comprises resin particles dispersed in a nonaq. solvent
AB
    having an elec. resistivity \geq 109~\Omega\cdot\text{cm} and a
     dielec. constant ≤3.5, wherein the resin particles have a
    multilayer structure and is made of a polymer which contains a unit
     [b1HCCb2(V0-L)] (V0 = COO, OCO, etc.; b1,2 = H, halo, cyano, etc.; and L =
    C8-32 alkyl, alkenyl) and a monofunctional monomer soluble in the solvent but
    becoming insol. upon polymerization and is obtained by seed polymerization in
the
    presence of seed grains having an average grain diameter 0.05-1.0 \mu m.
    liquid developer made the development-fixing processes faster and exhibited
     excellent development property for an electrophotog. plate making system
    using a large master plate.
ST
    liq developer dispersion resin particle electrophotog printing
    platemaking; seed polymn polymer liq developer dispersion resin particle
ΙT
    Polyesters, preparation
    RL: SPN (Synthetic preparation); TEM (Technical or engineered material
    use); PREP (Preparation); USES (Uses)
        (dispersion resin particle; liquid developer containing dispersion resin
        particles for electrophotog. printing platemaking system)
ΙT
    Printing plates
        (liquid developer containing dispersion resin particles for electrophotog.
       printing platemaking system)
    Electrophotographic developers
IT
        (liquid; liquid developer containing dispersion resin particles for
        electrophotog. printing platemaking system)
IT
     Polymerization
        (seed; liquid developer containing dispersion resin particles for
       electrophotog. printing platemaking system)
    85533-57-9P, Hexadecyl methacrylate-vinyl acetate copolymer
IT
                                                                   308283-85-4P
                                  473595-25-4P 473595-26-5DP, ester
                   473595-23-2P
     473595-22-1P
    473595-27-6DP, ester 473595-28-7DP, ester
    473595-29-8DP, ester 473595-30-1DP, ester
    473595-31-2DP, ester 473595-32-3DP, ester
    473595-33-4DP, ester 473595-34-5DP, ester
    473595-35-6DP, ester 473595-36-7DP, ester
                                                   473595-37-8DP, ester
    473595-38-9DP, ester 473595-40-3DP, ester
                                                   473595-41-4DP, ester
    473595-42-5DP, ester 473595-43-6DP, ester
                                                   473595-46-9DP, ester
    473595-44-7DP, ester 473595-45-8DP, ester
    473595-47-0DP, ester 473595-48-1DP, ester
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RL: SPN (Synthetic preparation); TEM (Technical or engineered material
     use); PREP (Preparation); USES (Uses)
        (dispersion resin particle; liquid developer containing dispersion resin
        particles for electrophotog. printing platemaking system)
     34888-27-2P, Dodecyl methacrylate-2-hydroxyethyl methacrylate copolymer
ΙT
     RL: SPN (Synthetic preparation); TEM (Technical or engineered material
     use); PREP (Preparation); USES (Uses)
        (dispersion stabilizing resin; liquid developer containing dispersion resin
        particles for electrophotog. printing platemaking system)
     473595-24-3DP, ester
IT
     RL: SPN (Synthetic preparation); TEM (Technical or engineered material
     use); PREP (Preparation); USES (Uses)
        (liquid developer containing dispersion resin particles for electrophotog.
        printing platemaking system)
     25053-53-6, Nucrel N699
IT
     RL: TEM (Technical or engineered material use); USES (Uses)
        (seed grain; liquid developer containing dispersion resin particles for
        electrophotog. printing platemaking system)
     9003-20-7P, Polyvinyl acetate
ΙT
     RL: SPN (Synthetic preparation); TEM (Technical or engineered material
     use); PREP (Preparation); USES (Uses)
        (seed polymer; liquid developer containing dispersion resin particles for
        electrophotog. printing platemaking system)
     473595-26-5DP, ester 473595-27-6DP, ester
IT
     473595-28-7DP, ester 473595-29-8DP, ester
     473595-30-1DP, ester 473595-31-2DP, ester
     473595-32-3DP, ester 473595-33-4DP, ester
     473595-34-5DP, ester 473595-42-5DP, ester
     473595-43-6DP, ester
     RL: SPN (Synthetic preparation); TEM (Technical or engineered material
     use); PREP (Preparation); USES (Uses)
        (dispersion resin particle; liquid developer containing dispersion resin
        particles for electrophotog. printing platemaking system)
     473595-26-5 HCAPLUS
RN
     2-Propenoic acid, 2-methyl-, 2-(diethylamino)ethyl ester, polymer with
CN
     dodecyl 2-methyl-2-propenoate, 2-hydroxyethyl 2-methyl-2-propenoate,
     methyl 2-methyl-2-propenoate, methyl 2-propenoate, octadecyl
     2-methyl-2-propenoate and 2-(phosphonooxy)ethyl 2-propenoate (9CI)
                                                                           (CA
     INDEX NAME)
     CM
          1
     CRN 32360-05-7
     CMF C22 H42 O2
               O CH<sub>2</sub>
Me^{-}(CH_2)_{17} - O^{-}C^{-}C^{-}Me
```

CM

CRN 32120-16-4 CMF C5 H9 O6 P

CM 3

CRN 868-77-9 CMF C6 H10 O3

CM 4

CRN 142-90-5 CMF C16 H30 O2

$$$^{\rm O}_{\rm H2}$$$
 Me— (CH2) 11 - O- C- C- Me

CM 5

CRN 105-16-8 CMF C10 H19 N O2

CM 6

CRN 96-33-3 CMF C4 H6 O2

$$\begin{array}{c} \text{O} \\ || \\ \text{MeO-C-CH----} \text{CH}_2 \end{array}$$

CM 7

CRN 80-62-6 CMF C5 H8 O2

RN 473595-27-6 HCAPLUS

CN 2-Butenedioic acid, decyl 2-[(1-oxo-2-propenyl)oxy]ethyl ester, polymer with 2-(diethylamino)ethyl 2-methyl-2-propenoate, methyl 2-methyl-2-propenoate, methyl 2-propenoate, 2-(phosphonooxy)ethyl 2-propenoate and tridecyl 2-methyl-2-propenoate (9CI) (CA INDEX NAME)

CM 1

CRN 305814-46-4 CMF C19 H30 O6

CM 2

CRN 32120-16-4 CMF C5 H9 O6 P

CM 3

CRN 2495-25-2 CMF C17 H32 O2

CM 4

CRN 105-16-8 CMF C10 H19 N O2

 \downarrow

CM 5

CRN 96-33-3 CMF C4 H6 O2

CM 6

CRN 80-62-6 CMF C5 H8 O2

RN 473595-28-7 HCAPLUS

CN Butanedioic acid, 1-[(2-methyl-1-oxo-2-propenyl)oxy]-3-[(1-oxohexyl)oxy]propyl pentyl ester, polymer with 2-(diethylamino)ethyl 2-methyl-2-propenoate, methyl 2-methyl-2-propenoate, methyl 2-propenoate, 2-(phosphonooxy)ethyl 2-propenoate and tridecyl 2-methyl-2-propenoate (9CI) (CA INDEX NAME)

CM 1

CRN 305814-48-6 CMF C22 H36 O8

CM 2

CRN 32120-16-4 CMF C5 H9 O6 P

CM 3

CRN 2495-25-2 CMF C17 H32 O2

CM 4

F

CRN 105-16-8 CMF C10 H19 N O2

CM 5

CRN 96-33-3 CMF C4 H6 O2

CM 6

CRN 80-62-6 CMF C5 H8 O2

RN 473595-29-8 HCAPLUS

CN Butanedioic acid, hexyl 3-[(2-methyl-1-oxo-2-propenyl)oxy]-2-[(1-oxoheptyl)oxy]propyl ester, polymer with 2-(diethylamino)ethyl 2-methyl-2-propenoate, methyl 2-methyl-2-propenoate, methyl 2-propenoate, 2-(phosphonoxy)ethyl 2-propenoate and tridecyl 2-methyl-2-propenoate (9CI) (CA INDEX NAME)

CM 1

CRN 305814-50-0 CMF C24 H40 O8 .

CM 2

CRN 32120-16-4 CMF C5 H9 O6 P

$$\begin{array}{c} \text{O} \\ \text{H}_2\text{O}_3\text{PO}-\text{CH}_2-\text{CH}_2-\text{O}-\text{C}-\text{CH} \Longrightarrow \text{CH}_2 \end{array}$$

CM3

CRN 2495-25-2 CMF C17 H32 O2

Me- (CH₂)
$$_{12}$$
 - O- C- C- Me

CM4

CRN 105-16-8 C10 H19 N O2 CMF

$$\begin{array}{c|c} {\rm H_2C} & {\rm O} \\ & || & || \\ {\rm Me-C-C-O-CH_2-CH_2-NEt_2} \end{array}$$

CM 5

CRN 96-33-3 CMF C4 H6 O2

CM 6

CRN 80-62-6 CMF C5 H8 O2

RN 473595-30-1 HCAPLUS

CN Nonanoic acid, 2-[[2-[(1-oxoheptyl)oxy]-3-[(1-oxo-2-propenyl)oxy]propyl]amino]ethyl ester, polymer with 2-(diethylamino)ethyl 2-methyl-2-propenoate, methyl 2-methyl-2-propenoate, methyl 2-propenoate, 2-(phosphonooxy)ethyl 2-propenoate and tridecyl 2-methyl-2-propenoate (9CI) (CA INDEX NAME)

CM 1

CRN 305814-52-2 CMF C24 H43 N O6

CM 2

CRN 32120-16-4 CMF C5 H9 O6 P

CM 3

CRN 2495-25-2 CMF C17 H32 O2

CM 4

CRN 105-16-8 CMF C10 H19 N O2

CM 5

CRN 96-33-3 CMF C4 H6 O2

CM 6

CRN 80-62-6 CMF C5 H8 O2

RN 473595-31-2 HCAPLUS

CN 2-Propenoic acid, 2-methyl-, 2-(diethylamino)ethyl ester, polymer with methyl 2-methyl-2-propenoate, methyl 2-propenoate, 2-(phosphonooxy)ethyl 2-propenoate, tridecyl 2-methyl-2-propenoate and 2-[[(tridecyloxy)carbonyl]amino]ethyl 2-methyl-2-propenoate (9CI) (CA INDEX NAME)

CM 1

CRN 305814-54-4 CMF C20 H37 N O4

CM 2

CRN 32120-16-4 CMF C5 H9 O6 P

$$^{\rm O}_{\rm H_2O_3PO^-CH_2^-CH_2^-O^-C^-CH^-CH_2^-CH_2^-}$$

CM 3

CRN 2495-25-2 CMF C17 H32 O2

$$^{\rm O}$$
 CH2 $_{||}$ $||$ Me— (CH2) 12 $^{\rm -}$ O— C— C— Me

CM 4

CRN 105-16-8 CMF C10 H19 N O2

CM 5

CRN 96-33-3 CMF C4 H6 O2

CM 6

CRN 80-62-6 CMF C5 H8 O2

RN 473595-32-3 HCAPLUS

CN 2-Propenoic acid, 2-methyl-, 2-(diethylamino)ethyl ester, polymer with dodecyl 2-propenoate, methyl 2-methyl-2-propenoate, methyl 2-propenoate, 2-(phosphonooxy)ethyl 2-propenoate and tridecyl 2-methyl-2-propenoate (9CI) (CA INDEX NAME)

$$Me^{-(CH_2)_{11}-O-C-CH} = CH_2$$

CRN 80-62-6 C5 H8 O2 CMF

RN473595-33-4 HCAPLUS

CN Decanoic acid, 2-[(2-methyl-1-oxo-2-propenyl)oxy]ethyl ester, polymer with 2-(diethylamino)ethyl 2-methyl-2-propenoate, methyl 2-methyl-2-propenoate, methyl 2-propenoate, 2-(phosphonooxy)ethyl 2-propenoate and tridecyl 2-methyl-2-propenoate (9CI) (CA INDEX NAME)

1 CM

CRN 32120-16-4 CMF C5 H9 O6 P

$$^{\circ}_{\text{H}_2\text{O}_3\text{PO}-\text{CH}_2-\text{CH}_2-\text{O}-\text{C}-\text{CH}}\!\!=\!\!\text{CH}_2$$

CM2

CRN 14792-62-2 CMF C16 H28 O4

CM 3

CRN 2495-25-2 CMF C17 H32 O2

$$$^{\rm O}_{\rm CH_2}$$$
 Me- (CH₂) $_{12}$ - O- C- C- Me

CM 4

CRN 105-16-8 CMF C10 H19 N O2

CRN 96-33-3 CMF C4 H6 O2

CM 6

CRN 80-62-6 CMF C5 H8 O2

RN 473595-34-5 HCAPLUS

CN Benzoic acid, 4-ethenyl-, octadecyl ester, polymer with 2-(diethylamino)ethyl 2-methyl-2-propenoate, methyl 2-methyl-2-propenoate, methyl 2-propenoate, 2-(phosphonooxy)ethyl 2-propenoate and tridecyl 2-methyl-2-propenoate (9CI) (CA INDEX NAME)

CM 1

CRN 308338-67-2 CMF C27 H44 O2

CM 2

CRN 32120-16-4 CMF C5 H9 O6 P

$$\begin{array}{c} \text{O} \\ || \\ \text{H}_2\text{O}_3\text{PO}-\text{CH}_2-\text{CH}_2-\text{O}-\text{C}-\text{CH} \end{array} \text{CH}_2$$

CRN 2495-25-2 CMF C17 H32 O2

CM4

CRN 105-16-8 C10 H19 N O2 CMF

CM5

CRN 96-33-3 C4 H6 O2 CMF

CM 6

CRN 80-62-6 CMF C5 H8 O2

473595-42-5 HCAPLUS RN

2-Propenoic acid, 2-methyl-, 2-(diethylamino)ethyl ester, polymer with CN 4-ethenylphenyl dihydrogen phosphate, methyl 2-methyl-2-propenoate, methyl 2-propenoate, octadecyl 2-methyl-2-propenoate and tridecyl 2-methyl-2-propenoate (9CI) (CA INDEX NAME)

CRN 80122-59-4 CMF C8 H9 O4 P

CM 2

CRN 32360-05-7 CMF C22 H42 O2

CM 3

CRN 2495-25-2 CMF C17 H32 O2

CM 4

CRN 105-16-8 CMF C10 H19 N O2

CM 5

CRN 96-33-3 CMF C4 H6 O2

CRN 80-62-6 CMF C5 H8 O2

RN 473595-43-6 HCAPLUS

CN 2-Propenoic acid, 2-methyl-, 2-(diethylamino)ethyl ester, polymer with methyl 2-methyl-2-propenoate, methyl 2-propenoate, octadecyl 2-methyl-2-propenoate, [4-[(1-oxo-2-propenyl)amino]phenyl]phosphonic acid and tridecyl 2-methyl-2-propenoate (9CI) (CA INDEX NAME)

-CM 1

CRN 149234-87-7 CMF C9 H10 N O4 P

CM 2

CRN 32360-05-7 CMF C22 H42 O2

CM 3

CRN 2495-25-2 CMF C17 H32 O2

$$$^{\rm O}$$$
 CH2 $$^{\rm H}_{\rm 2}$$ Me- (CH2)12-O-C-C-Me

CRN 105-16-8 CMF C10 H19 N O2

CM5

CRN 96-33-3 CMF C4 H6 O2

CM6

CRN 80-62-6 CMF C5 H8 O2

L23 ANSWER 5 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 2002:748408 HCAPLUS

137:302107 DN

ED Entered STN: 03 Oct 2002

Electrostatographic liquid developers for making electrographic printing ΤI plates

Kato, Eiichi ΙN

Fuji Photo Film Co., Ltd., Japan PA

Jpn. Kokai Tokkyo Koho, 33 pp. SO CODEN: JKXXAF

 DT Patent

Japanese LA

ICM G03G009-13 IC ICS C08F291-00

CC 74-3 (Radiation Chemistry, Photochemistry, and Photographic and Other Reprographic Processes)

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Section cross-reference(s): 35
FAN.CNT 1
     PATENT NO.
                      KIND DATE
                                           APPLICATION NO. DATE
                    ____
                                           JP 2001-85195 20010323
    JP 2002287434 A2 20021003
PΙ
PRAI JP 2001-85195
                            20010323
     The title developer contains dispersed resin particles, which are prepared
     from monomers in an aprotic solvent in the present of dispersion-
     stabilizing resin, in an aprotic solvent of \geq 109~\Omega\cdot\text{cm}
     and of ≤3.5 dielec. constant, wherein the dispersed resin
     particles are made of monomers: a monomer having a functional group, which
     is soluble in an aprotic solvent and becomes insol. in the solvent after the
     polymerization; a monomer having -N(R1)(R2) group ( R1-2 = H, C1-22
hydrocarbon);
     and a monomer having an acidic group chosen from -PO3H2, -SO3H, and -SO2H,
     wherein the dispersion stabilizing resin is made of a star burst polymer,
     which has ≥3 A-B block polymer chains connected to a core organic
     group and 2X104-1X106 weight average mol. weight The block A is made of a
monomer
     having a functional group, which is soluble in an aprotic solvent and becomes
     insol. in the solvent after the polymerization and a monomer having polar group
     chosen from phosphono, carboxyl, sulfo, hydroxyl, formyl, amino,
     -P(=0)(OH)E1 (E1 = hydrocarbon, oxyhydrocarbon), and cyclic acid
     anhydride. The block B has [-CH(b1)-C(B2)(A-L)]-(A = -COO-,
     -(CH2)\times COO-, -(CH2)\times OCO- ( \times = 1-3 integer), etc. L =
     C \ge 8 aliphatics, B1-2 = H, halo, cyano, C1-7 hydrocarbon, etc.). The
     developer provides the rapid development/fixing process for large
     electrophotog. printing plate masters and shows the good characteristics
     as a liquid developers.
\mathtt{ST}
     electrostatog liq developer electrog printing plate
     Printing plates
\operatorname{IT}
        (electrog.; electrostatog. liquid developers for making electrog.
        printing plates)
ΙT
     Electrographic developers
        (electrostatog. liquid developers for making electrog. printing plates)
     Dendritic polymers
IT
     RL: SPN (Synthetic preparation); TEM (Technical or engineered material
     use); PREP (Preparation); USES (Uses)
        (electrostatog. liquid developers for making electrog. printing plates)
                   150551-89-6 150551-92-1 150551-93-2 150551-97-6
IT
     150551-83-0
                   155293-25-7 159967-38-1 159967-39-2
     154340-06-4
                                                             159967-40-5
     159967-41-6 159967-42-7 159967-43-8 159967-44-9
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (core; electrostatog. liquid developers for making electrog. printing
       plates)
                   159967-45-0, Styrene-4-methylstyrene-octadecyl
    159967-35-8
\operatorname{IT}
    methacrylate-dodecyl acrylate block copolymer 159967-46-1 159967-47-2
     159967-48-3 159967-49-4 159967-50-7 159967-51-8
                                                             159967-52-9
                 159967-54-1 159967-55-2
     159967-53-0
                                              159967-56-3,
     Styrene-4-hydroxystyrene-tetradecyl methacrylate block copolymer
     467435-80-9
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (dendritic; electrostatog. liquid developers for making electrog.
        printing plates)
    150469-20-8P, Methyl methacrylate-methyl acrylate-stearyl acrylate block
IT
     copolymer
    RL: SPN (Synthetic preparation); TEM (Technical or engineered material
    use); PREP (Preparation); USES (Uses)
        (dendritic; electrostatog. liquid developers for making electrog.
```

printing plates)

IT 9003-20-7, Vinyl acetate homopolymer

RL: RCT (Reactant); RACT (Reactant or reagent)

(resin in developer; electrostatog. liquid developers for making electrog. printing plates)

26246-92-4P, Dodecyl acrylate homopolymer 29500-86-5P, Decyl acrylate IThomopolymer 41630-11-9P, Tridecyl methacrylate homopolymer 80122-60-7P 442156-57-2P, Methyl methacrylate-methyl 329914-76-3P acrylate-2-(N, N-dimethylamino) ethyl methacrylate-octadecyl acrylate-2-phosphonoethyl methacrylate copolymer 467435-84-3DP, Me methacrylate/ethyl acrylate/2-(N, N-diethylamino)ethyl acrylate/2,3-dioctanoyloxypropyl methacrylate/4-phosphobutyl acrylate **copolymer** 467435-85-4P 467435-86-5P 467435-88-7P 467435-90-1P 467435-92-3P 467435-91-2P 467435-93-4P 467435-95-6P 467435-98-9P 467436-00-6P 467436-02-8P 467435-96-7P RL: SPN (Synthetic preparation); TEM (Technical or engineered material

use); PREP (Preparation); USES (Uses) (resin in developer; electrostatog. liquid developers for making electrog. printing plates)

IT 467435-84-3DP, Me methacrylate/ethyl acrylate/2-(N,N-diethylamino)ethyl acrylate/2,3-dioctanoyloxypropyl methacrylate/4-phosphobutyl acrylate copolymer

RL: SPN (Synthetic preparation); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses)

(resin in developer; electrostatog. liquid developers for making electrog. printing plates)

RN 467435-84-3 HCAPLUS

CN Octanoic acid, 1-[[(2-methyl-1-oxo-2-propenyl)oxy]methyl]-1,2-ethanediyl ester, polymer with 2-(diethylamino)ethyl 2-propenoate, methyl 2-methyl-2-propenoate, methyl 2-propenoate and 4-(phosphonooxy)butyl 2-propenoate (9CI) (CA INDEX NAME)

CM 1

CRN 154732-35-1 CMF C23 H40 O6

$$H_{2}C$$
 O $||$ || || Me-C-C-O-CH₂ O $||$ || Me-(CH₂)6-C-O-CH₂-CH-O-C-(CH₂)6-Me

CM 2

CRN 110507-31-8 CMF C7 H13 O6 P

$$^{\circ}_{\text{H}_2\text{O}_3\text{PO}}$$
 (CH₂)₄-0-C-CH== CH₂

CRN 2426-54-2 CMF C9 H17 N O2

$$\begin{array}{c} \text{O} \\ \parallel \\ \text{Et}_2 \text{N} - \text{CH}_2 - \text{CH}_2 - \text{O} - \text{C} - \text{CH} \Longrightarrow \text{CH}_2 \end{array}$$

CM

CRN 96-33-3 C4 H6 O2 CMF

CM5

CRN 80-62-6 C5 H8 O2 CMF

L23 ANSWER 6 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

2002:638105 HCAPLUS AN

DN137:181915

Entered STN: 23 Aug 2002 ED

Entered STN: 23 Aug 2002
Phosphorus-containing polymers for optical signal transducers TIapplicante

IN Dorn, Ingmar; Kohler, Burkhard

Germany PA

U.S. Pat. Appl. Publ., 12 pp. SO CODEN: USXXCO

 DT Patent

English LA

ICM G02B006-22 IC ICS C08K005-49

NCL 385128000

9-2 (Biochemical Methods) CC

Section cross-reference(s): 3, 7, 15, 38, 79, 80

FAN.	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 2002114604	A1	20020822	US_2002-81628	20020220
	DE 10108483	A1	20020905	DE 2001-10108483	20010222
	WO 2002068481	A1	20020906	WO 2002-EP1399	20020211

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W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
             PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
             UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU,
             TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,
             CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
             BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                            20031203
                       Α1
                                           EP 2002-704708
     EP 1366088
                                                            20020211
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
                            20010222
PRAI DE 2001-10108483
                       Α
     WO 2002-EP1399
                            20020211
     Phosphorus-containing polymers suitable for coating dielec. surfaces
AB
     are described by the general formulas P(A)m(F)n1(U)o1 (I) and
     P(A)m(UFn2)o2 (II) (P = (un)branched, (un)crosslinked homo- or
     heteropolymeric polymer component; A = identical or different
     phosphorus-containing groups bonded to P; m = .apprx.3-1000, F = identical or
     different functional groups bonded directly or indirectly to P; n1 =
     .apprx.1-1000; n2 = .apprx.1-100, U = identical or different (un)branched
     (un)crosslinked oligomeric or polymeric segments made up of identical or
     different monomers which are bonded to P; o1 = .apprx.0-1000, o2 =
     .apprx.1-1000). Methods for preparing the polymers are described which
     entail copolymg. a monomer containing a phosphorus-containing group A, or a
     plurality of identical or different monomers containing identical or different
     phosphorus-containing groups A, with a monomer containing a functional group
F, or
     a plurality of identical or different monomers containing identical or
     different functional groups F, and optionally, a monomer containing a segment
     U, or a plurality of identical or different monomers containing identical or
     different segments U, to form I, or with a monomer containing a unit (UFn2)o2,
     or a plurality of identical or different monomers containing identical or
     different units of the formula (UFn2)o2, to form II. The use of the
     polymers for coating dielec. materials, in particular
     dielec. waveguides, and optical signal
     transducers with dieled. waveguides coated by the
     polymers are also described. The optical signal transducers
     having a coated dielec. waveguides may be used for
     immobilizing chemical and/or biochem. recognition elements.
     phosphorus contg polymer prepn; dielec waveguide
ST
     coating phosphorus contg polymer; optical signal transducer
     phosphorus contg polymer coated dielec waveguide
     Antibodies and Immunoglobulins
IT
     RL: ARG (Analytical reagent use); DEV (Device component use); ANST
     (Analytical study); USES (Uses)
        (immobilized; phosphorus-containing polymers and their preparation and
their use
        for coating dielec. waveguides and optical signal
        transducers using the coated waveguides)
IT
    Biosensors
        (immunosensors; phosphorus-containing polymers and their preparation
        and their use for coating dielec. waveguides and
        optical signal transducers using the coated
        wavequides)
IT
    Biosensors
        (optical; phosphorus-containing polymers and their preparation and their
use for
```

coating dielec. waveguides and optical signal transducers using the coated waveguides) Immobilization, molecular or cellular ITOptical sensors Wavequides (phosphorus-containing polymers and their preparation and their use for coating dielec. waveguides and optical signal transducers using the coated waveguides) Polyphosphoric acids IT RL: IMF (Industrial manufacture); PEP (Physical, engineering or chemical process); PYP (Physical process); TEM (Technical or engineered material use); PREP (Preparation); PROC (Process); USES (Uses) (reaction products with polymers; phosphorus-containing polymers and their preparation and their use for coating dielec. waveguides and optical signal transducers using the coated waveguides) Fatty acids, preparation ITRL: IMF (Industrial manufacture); PEP (Physical, engineering or chemical process); PYP (Physical process); TEM (Technical or engineered material use); PREP (Preparation); PROC (Process); USES (Uses) (soya, reaction products with glycidol and phosphoric acid; phosphorus-containing polymers and their preparation and their use for coating dielec. waveguides and optical signal transducers using the coated waveguides) 1314-23-4, Zirconium oxide, uses 1314-61-0, Tantalum oxide 1344-28-1, ITAluminum oxide, uses 12055-23-1, Hafnium oxide 13463-67-7, Titanium oxide, uses RL: DEV (Device component use); PEP (Physical, engineering or chemical process); PYP (Physical process); PROC (Process); USES (Uses) (phosphorus-containing polymers and their preparation and their use for coating dielec. waveguides and optical signal transducers using the coated waveguides) 108-31-6DP, Maleic Acid Anhydride, reaction products with polyglycidols IT556-52-5DP, Glycidol, reaction products with fatty acids and phosphoric acid 1066-51-9DP, Aminomethanephosphonic acid, reaction products with polymers 1746-03-8DP, Vinylphosphonic acid, reaction products with polyglycidols 7664-38-2DP, Phosphoric acid, reaction products with polylysine salts 7664-38-2DP, Phosphoric acid, reaction products with polymers 9002-89-5DP, Polyvinyl alcohol, reaction products with polyphosphoric acid 9011-16-9DP, Maleic anhydride-methyl vinyl ether copolymer, reaction products with aminomethanephosphonic acid 9041-77-4P, Dextran phosphate 17261-34-6DP, Iminobismethylene Phosphonic Acid, reaction products with polyglycidols 21282-97-3DP, reaction products with polyglycidols and vinylphosphonic acid 25988-63-0DP, Poly-L-lysine hydrobromide, reaction products with phosphoric acid 26588-20-5DP, reaction products with phosphoric acid 69680-04-2DP, reaction products with phosphoric 98980-94-0DP, reaction products with iminobismethylene phosphonic acid and maleic acid anhydride 449188-13-0P RL: IMF (Industrial manufacture); PEP (Physical, engineering or chemical process); PYP (Physical process); TEM (Technical or engineered material use); PREP (Preparation); PROC (Process); USES (Uses) (phosphorus-containing polymers and their preparation and their use for coating dielec. waveguides and optical signal transducers using the coated waveguides) 156-57-0 13598-36-2, Phosphonic acid IT

RL: RCT (Reactant); RACT (Reactant or reagent) (phosphorus-containing polymers and their preparation and their use for coating dielec. waveguides and optical signal transducers using the coated waveguides) 1066-51-9DP, Aminomethanephosphonic acid, reaction products with IT polymers 1746-03-8DP, Vinylphosphonic acid, reaction products with polyglycidols 7664-38-2DP, Phosphoric acid, reaction products with polylysine salts 17261-34-6DP, Iminobismethylene Phosphonic Acid, reaction products with polyglycidols 449188-13-0P RL: IMF (Industrial manufacture); PEP (Physical, engineering or chemical process); PYP (Physical process); TEM (Technical or engineered material use); PREP (Preparation); PROC (Process); USES (Uses) (phosphorus-containing polymers and their preparation and their use for coating dielec. waveguides and optical signal transducers using the coated waveguides) RN1066-51-9 HCAPLUS Phosphonic acid, (aminomethyl) - (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME) CNH2N-CH2-PO3H2 RN 1746-03-8 HCAPLUS Phosphonic acid, ethenyl- (9CI) (CA INDEX NAME) CN. $H_2C = CH - PO_3H_2$ 7664-38-2 HCAPLUS RN Phosphoric acid (7CI, 8CI, 9CI) (CA INDEX NAME) CNOH RN 17261-34-6 HCAPLUS Phosphonic acid, [iminobis(methylene)]bis- (9CI) (CA INDEX NAME) CN H2O3P-CH2-NH-CH2-PO3H2 RN449188-13-0 HCAPLUS Butanoic acid, 3-oxo-, 2-[(2-methyl-1-oxo-2-propenyl)oxy]ethyl ester, CN polymer with ethenylphosphonic acid and α -(1-oxo-2-propenyl)- ω methoxypoly(oxy-1,2-ethanediyl) (9CI) (CA INDEX NAME) CMCRN 32171-39-4 CMF (C2 H4 O)n C4 H6 O2 CCI PMS

$$H_2C = CH - C - CH_2 - CH_2 - CH_2 - OMe$$

CRN 21282-97-3 CMF C10 H14 O5

CM 3

CRN 1746-03-8 CMF C2 H5 O3 P

 $H_2C = CH - PO_3H_2$

L23 ANSWER 7 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

2001:371617 HCAPLUS AN

135:6996 DN

Entered STN: 23 May 2001 ED

Oil-based inks with good deliverability and image-forming properties for TIelectrostatic ink-jet printing

Kato, Eiichi IN

PAFuji Photo Film Co., Ltd., Japan

SO Jpn. Kokai Tokkyo Koho, 48 pp.

CODEN: JKXXAF

 DT Patent

LA Japanese

ICM C09D011-00 IC

ICS B41J002-01; B41M005-00

CC 42-12 (Coatings, Inks, and Related Products) Section cross-reference(s): 74

FAN.CNT 1

APPLICATION NO. PATENT NO. KIND DATE DATE 20010522 JP 2000-261060 JP 2001139860 A2 ΡI 20000830 PRAI JP 1999-246120 A 19990831

The inks dispersed in a nonaq. medium having elec. resistance ≥109 AB Ω -cm and **dielec**. constant ≤ 3.5 contain resin particles manufactured by polymerizing solns. containing (A) ≥ 1 nonaq. solvent-soluble monofunctional monomers which become insol. in the nonag. solvents after being polymerized, (B) ≥ 1 macromonomers (Mw ≤ 2 x 104) having repeating units containing fluoro and/or silyl groups and terminated at one end with polymerizable double bond, and (C) ≥ 1

partially crosslinked and nonaq. solvent-soluble polymeric dispersion stabilizers. Thus, vinyl acetate was polymerized with Silaplane FM 0721 (methacrylate- and trimethylsilyl-terminated polydimethylsiloxane) in the presence of octadecyl methacrylate-divinylbenzene copolymer in Isopar H (isoalkanes) and filtered to give particles, which was dispersed with alkali blue dispersion in Isopar E (isoalkane) to give an ink. oil based ink polysiloxane graft deliverability; jet printing ink ST electrostatic polysiloxane dispersion; vinylbenzene octadecyl methacrylate dispersant polysiloxane ink; vinyl acetate polydimethylsiloxane macromonomer graft ink Polysiloxanes, uses ITRL: IMF (Industrial manufacture); PRP (Properties); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses) (graft polymers, acrylic; oil-based inks with good deliverability and image-forming properties for electrostatic ink-jet printing) ΙT Inks (jet-printing; oil-based inks with good deliverability and image-forming properties for electrostatic ink-jet printing) ITInks (lithog.; oil-based inks with good deliverability and image-forming properties for electrostatic ink-jet printing) ITTelomers (polymers) RL: IMF (Industrial manufacture); POF (Polymer in formulation); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses) (oil-based inks with good deliverability and image-forming properties for electrostatic ink-jet printing) IT Inks (oil-based; oil-based inks with good deliverability and image-forming properties for electrostatic ink-jet printing) ITDispersing agents (reactive, macromer; oil-based inks with good deliverability and image-forming properties for electrostatic ink-jet printing) 61255-17-2P, Divinylbenzene-dodecyl methacrylate copolymer IT122324-74-7P, Divinylbenzene-octadecyl methacrylate copolymer 130805-26-4P, Divinylbenzene-hexadecyl methacrylate copolymer 148532-67-6P, Dodecyl methacrylate-octyl methacrylate-trivinylbenzene copolymer 148532-68-7P, Butyl methacrylate-ethylene glycol dimethacrylate-octadecyl methacrylate 148640-01-1P, Divinylbenzene-octadecyl methacrylatecopolymer thioglycolic acid telomer ester with 2-hydroxyethyl methacrylate 159446-39-6P, Divinylbenzene-octadecyl methacrylate-2-mercaptoethanol telomer ester with 10-carboxyldecylacrylamide 159446-41-0P 159446-42-1P, Divinylbenzene-octadecyl methacrylate-2-mercaptoethanol telomer ester with 4-vinylbenzenecarboxylic acid 159446-44-3P, Divinylbenzene-octadecyl methacrylate-2-mercaptoethanol telomer ester with vinylacetic acid 159446-45-4P, Divinylbenzene-octadecyl methacrylate-2-mercaptoethanol telomer ester with methacrylic acid 159446-48-7P, Divinylbenzene-octadecyl methacrylate-2-mercaptoethanol telomer ester with acrylic acid 214772-24-4P, Divinylbenzene-octadecyl methacrylate-2-mercaptoethanol telomer ester with 2-carboxyethyl acrylate 214772-26-6P, Divinylbenzene-octadecyl methacrylate-2-mercaptoethanol telomer ester with α -chloroacrylic acid 214772-29-9P, Divinylbenzene-octadecyl methacrylate-2-mercaptoethanol telomer ester with 2-(2-carboxyethylcarbonyloxy)ethyl cyanoacrylate 218459-53-1P, Allyl methacrylate-dodecyl methacrylate-thioglycolic acid telomer ester with 2-hydroxyethyl methacrylate 218459-59-7P, Ethylene glycol dimethacrylate-octadecyl methacrylate-thioglycolic acid telomer ester with 2-hydroxyethyl methacrylate 218459-61-1P, Hexadecyl methacrylatepropylene glycol dimethacrylate-thioglycolic acid telomer ester with 218459-65-5P, Butyl methacrylate-divinyl 2-hydroxyethyl methacrylate

IT

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adipate-dodecyl methacrylate-thioglycolic acid telomer ester with
   2-hydroxyethyl methacrylate
                                218459-67-7P, Ethylene glycol
   diacrylate-methyl methacrylate-octadecyl methacrylate-thioglycolic acid
   telomer ester with 2-hydroxyethyl methacrylate
                                                   218459-72-4P,
   Divinylbenzene-styrene-tetradecyl methacrylate-thioglycolic acid telomer
   ester with 2-hydroxyethyl methacrylate 324529-94-4P, Ethylene glycol
   diacrylate-hexadecyl methacrylate copolymer 324574-61-0P
                                                               341506-19-2P
   RL: IMF (Industrial manufacture); MOA (Modifier or additive use); PREP
   (Preparation); USES (Uses)
      (dispersant; oil-based inks with good deliverability and image-forming
      properties for electrostatic ink-jet printing)
   139703-31-4P, Divinylbenzene-Octadecyl methacrylate-thioglycolic acid
            139703-33-6P, Divinylbenzene-tridecyl methacrylate-thioglycolic
   telomer
   acid telomer
                  139720-57-3P, Divinylbenzene-Octadecyl methacrylate-3-
   thiopropionic acid telomer
                               139720-59-5P
                                              139720-60-8P
   139720-62-0P
                                139720-64-2P, Octadecyl methacrylate-
                  139720-63-1P
   divinylbenzene-2-mercaptoethylamine telomer 141181-86-4P,
   Divinylbenzene-dodecyl methacrylate-thioglycolic acid telomer
   148532-76-7P, Octadecyl methacrylate-butyl methacrylate-ethylene glycol
   dimethacrylate-thioglycolic acid telomer 148532-82-5P, Hexadecyl
   methacrylate-divinyl adipate-thioglycolic acid telomer
                                                           159291-22-2P,
   Trivinylbenzene-dodecyl methacrylate-octyl methacrylate-thioglycolic acid
   telomer
             159291-24-4P
                           215672-71-2P
                                          308283-76-3P, Docosyl
   methacrylate-polyethylene glycol diacrylate copolymer
   RL: IMF (Industrial manufacture); MOA (Modifier or additive use); PREP
   (Preparation); USES (Uses)
      (oil-based inks with good deliverability and image-forming properties
      for electrostatic ink-jet printing)
   80-62-6DP, polymers with (meth)acrylates
                                             96-33-3DP, polymers with
                    106-91-2DP, polymers with (meth) acrylates 2867-47-2DP,
   (meth)acrylates
   polymers with (meth)acrylates 7582-21-0DP, polymers with (meth)acrylates
   80730-17-2DP, polymers with (meth)acrylates
   152792-47-7DP, polymers with (meth)acrylates
   169045-89-0P 305814-07-7DP, polymers with
   (meth) acrylates 305814-10-2DP, polymers with
   (meth)acrylates
                    308278-98-0DP, polymers with (meth)acrylates
   311807-05-3DP, polymers with (meth)acrylates
                                                 311807-06-4P, Silaplane FM
   0721-vinyl acetate graft copolymer 340756-70-9DP, polymers with
                                   341031-31-0P
                    341031-29-6P
   (meth)acrylates
                                                  341031-32-1P
   341031-33-2P
                 341031-35-4P
                                341031-36-5P
                                               341031-38-7P
                                                              341031-39-8P
   341031-40-1P
                                341031-42-3P
                                               341031-43-4P
                 341031-41-2P
                                                              341031-44-5P
                341031-46-7P
341031-45-6P
                                341505-86-0P
                                               341505-91-7P
                                                              341505-93-9P
                                341505-96-2P 341505-98-4P
   341505-94-0P 341505-95-1P
                                                              341506-00-1P
   341506-01-2P 341506-30-7P
                                               341506-44-3P
                                341506-35-2P
                                                              341506-46-5P
   341506-51-2P 341506-56-7P
   RL: IMF (Industrial manufacture); PRP (Properties); TEM (Technical or
   engineered material use); PREP (Preparation); USES (Uses)
      (oil-based inks with good deliverability and image-forming properties
    for electrostatic ink-jet printing)
  80730-17-2DP, polymers with (meth)acrylates
   152792-47-7DP, polymers with (meth)acrylates
   305814-07-7DP, polymers with (meth)acrylates
   305814-10-2DP, polymers with (meth)acrylates
   RL: IMF (Industrial manufacture); PRP (Properties); TEM (Technical or
   engineered material use); PREP (Preparation); USES (Uses)
      (oil-based inks with good deliverability and image-forming properties
     for electrostatic ink-jet printing)
   80730-17-2 HCAPLUS
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2-Propenoic acid, 2-methyl-, 2-phosphonoethyl ester (9CI) (CA INDEX NAME)

RN 152792-47-7 HCAPLUS

CN Benzenemethanol, 4-ethenyl-, dihydrogen phosphate (9CI) (CA INDEX NAME)

RN 305814-07-7 HCAPLUS

CN 2-Propenoic acid, 2-methyl-, 5-phosphonopentyl ester (9CI) (CA INDEX NAME)

RN 305814-10-2 HCAPLUS

CN 2-Propenamide, N-[10-(phosphonooxy)decyl]- (9CI) (CA INDEX NAME)

L23 ANSWER 8 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 2000:658812 HCAPLUS

DN 134:53235

ED Entered STN: 20 Sep 2000

TI Glucose sensor with improved hemocompatibilty

AU Yang, Y.; Zhang, S. F.; Kingston, M. A.; Jones, G.; Wright, G.; Spencer, S. A.

CS Centre for Science and Technology in Medicine, Keele University, Staffordshire, ST5 5BG, UK

SO Biosensors & Bioelectronics (2000), 15(5-6), 221-227 CODEN: BBIOE4; ISSN: 0956-5663

PB Elsevier Science S.A.

DT Journal

LA English

CC 9-1 (Biochemical Methods)

AB A new biocompatible copolymer has been synthesized and used in an electrochem. enzyme-based glucose sensor. The copolymer incorporates three segments including a monomer with an elec. neutral phosphorylcholine head group that is able to reject protein adsorption and two segments that increase the affinity to polyurethane substrate. Peel and solution circulation tests showed that this material has high attachment

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to polyurethane. With the new copolymer as the outermost layer and the
     polyurethane as the diffusion-limiting membrane, the sensor
     showed extended linearity up to 50 mM glucose and stable output in bovine
     serum for 70 h. During in vivo tests, the sensor exhibited a
     steady current signal and a rapid transient response when the glucose
     concentration was raised. These results imply that the hemocompatibility of
the
     glucose sensor coated with the new copolymer has been improved,
     which is crucial for a sensor used for clin. real-time
     monitoring. The material may also be suitable for application to other
     implantable devices.
     glucose sensor hemocompatibilty copolymer
ST
     Biocompatibility
\operatorname{IT}
     Blood analysis
     Glucose sensors
     Temperature
        (glucose sensor with improved hemocompatibilty)
IT
     Polyurethanes, uses
     RL: DEV (Device component use); USES (Uses)
        (glucose sensor with improved hemocompatibilty)
     Adsorption
IT
        (protein; glucose sensor with improved hemocompatibilty)
     50-99-7, D-Glucose, analysis
IT
     RL: ANT (Analyte); ANST (Analytical study)
        (glucose sensor with improved hemocompatibilty)
ΙT
     125275-25-4DP, copolymer with another vinyl monomer, 3C-
     copolymer
     RL: DEV (Device component use); SPN (Synthetic preparation);
     PREP (Preparation); USES (Uses)
        (glucose sensor with improved hemocompatibilty)
ΙT
     78-67-1
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (glucose sensor with improved hemocompatibilty)
              THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
       18
RE
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(4) Clark, L; Ann New York Acad Sci 1962, V102, P29 HCAPLUS
(5) Ghosh, S; Abstracts of papers of the American Chemical Society 1994,
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    125275-25-4DP, copolymer with another vinyl monomer, 3C-
    copolymer
    RL: DEV (Device component use); SPN (Synthetic preparation);
    PREP (Preparation); USES (Uses)
        (glucose sensor with improved hemocompatibilty)
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RN 125275-25-4 HCAPLUS

CN 3,5,8-Trioxa-4-phosphaundec-10-en-1-aminium, 4-hydroxy-N,N,N,10-tetramethyl-9-oxo-, inner salt, 4-oxide, polymer with butyl 2-methyl-2-propenoate (9CI) (CA INDEX NAME)

CM 1

4.

CRN 67881-98-5 CMF C11 H22 N O6 P

CM 2

CRN 97-88-1 CMF C8 H14 O2

L23 ANSWER 9 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1999:563650 HCAPLUS

DN 132:134206

ED Entered STN: 06 Sep 1999

TI Multifunctional biocompatible membrane and its application to fabricate a miniaturized glucose **sensor** with potential for use in vivo

AU Chen, Chien-Yuan; Ishihara, Kazuhiko; Nakabayashi, Nobuo; Tamiya, Eiichi; Karube, Isao

CS Department of Agricultural Chemistry, National Taiwan University, Taipei, 106, Taiwan

SO Biomedical Microdevices (1999), 1(2), 155-166 CODEN: BMICFC; ISSN: 1387-2176

PB Kluwer Academic Publishers

DT Journal

LA English

CC **9-1** (**Biochemical** Methods)

AB A multifunctional membrane with biocompatibility, diffusion-limiting effect, and the ability to curtail the responses of an H2O2 electrode to ascorbate and urate was prepared. It was composed of MB, AB, and CTA, where MB is the copolymer of 2-methacryloyloxyethyl phosphorylcholine (MPC) and n-butyl-methacrylate (BMA), AB is the copolymer of acrylamide-2-methylpropane sulfonic acid (AMPS) and BMA, CTA is cellulose triacetate. Investigation of the biocompatibility of this membrane showed that, compared with CTA, relatively few platelets bound to it. The membrane was coated onto the working electrode of a needle-type glucose sensor on which immobilized glucose oxidase membrane has been coated. The sensor did not respond to ascorbate and urate at their concentration normally encountered in blood. Its response was not inhibited by metal

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ions in blood at usual concentration The sensor exhibited superior thermostability in addition to a rapid response (< 90 s in batch operation), good reproducibility (RE < 5%), good stability (more than 36 h continuously in heparinized whole blood), and a wide dynamic range (5-650 mg/dL glucose). The sensor was used to determine glucose in serum. The data obtained from the sensor showed good agreement with that from a clin. automated analyzer (R = 0.973). multifunctional biocompatible membrane miniaturized glucose sensor electrode Platelet (blood) (adhesion; multifunctional biocompatible membrane and its application to fabricate a miniaturized glucose sensor with potential for use in vivo) Membranes, nonbiological (biocompatible; multifunctional biocompatible membrane and its application to fabricate a miniaturized glucose sensor with potential for use in vivo) Cations (effect; multifunctional biocompatible membrane and its application to fabricate a miniaturized glucose sensor with potential for use in vivo) Blood analysis (glucose; multifunctional biocompatible membrane and its application to fabricate a miniaturized glucose sensor with potential for use in vivo) Biocompatibility Glucose sensors Thermal stability (multifunctional biocompatible membrane and its application to fabricate a miniaturized glucose sensor with potential for use in vivo) Cell adhesion (platelet; multifunctional biocompatible membrane and its application to fabricate a miniaturized glucose sensor with potential for use in vivo) 50-81-7, Ascorbic acid, analysis 69-93-2, Uric acid, analysis RL: ARU (Analytical role, unclassified); ANST (Analytical study) (interferent, no effect; multifunctional biocompatible membrane and its application to fabricate a miniaturized glucose sensor with potential for use in vivo) 50-99-7, D-Glucose, analysis RL: ANT (Analyte); BOC (Biological occurrence); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study); OCCU (Occurrence) (multifunctional biocompatible membrane and its application to fabricate a miniaturized glucose sensor with potential for use in vivo) 9001-37-0D, Glucose oxidase, immobilized RL: ARG (Analytical reagent use); DEV (Device component use); ANST (Analytical study); USES (Uses) (multifunctional biocompatible membrane and its application to fabricate a miniaturized glucose sensor with potential for use in vivo) 256494-85-6P RL: ARU (Analytical role, unclassified); DEV (Device component use); POF (Polymer in formulation); SPN (Synthetic preparation); ANST (Analytical study); PREP (Preparation); USES (Uses) (multifunctional biocompatible membrane and its application to fabricate a miniaturized glucose sensor with potential foruse in vivo)

IT 9012-09-3, Cellulose triacetate 64112-04-5 125275-25-4
RL: POF (Polymer in formulation); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(multifunctional biocompatible membrane and its application to fabricate a miniaturized glucose **sensor** with potential for use in vivo)

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IT **256494-85-6P**

RL: ARU (Analytical role, unclassified); DEV (Device component use); POF (Polymer in formulation); SPN (Synthetic preparation); ANST (Analytical study); PREP (Preparation); USES (Uses)

(multifunctional biocompatible membrane and its application to fabricate a miniaturized glucose sensor with potential for use in vivo)

RN 256494-85-6 HCAPLUS

CN 3,5,8-Trioxa-4-phosphaundec-10-en-1-aminium, 4-hydroxy-N,N,N,10tetramethyl-9-oxo-, inner salt, 4-oxide, polymer with butyl 2-methyl-2-propenoate, 2-methyl-2-[(1-oxo-2-propenyl)amino]-1propanesulfonic acid and 1,1,2,2-tetrachloroethane (9CI) (CA INDEX NAME)

CM 1

CRN 67881-98-5 CMF C11 H22 N O6 P

CM 2

CRN 15214-89-8 CMF C7 H13 N O4 S

CM 3

CRN 97-88-1 CMF C8 H14 O2

CRN 79-34-5 CMF C2 H2 C14

IT 125275-25-4

RL: POF (Polymer in formulation); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(multifunctional biocompatible membrane and its application to fabricate a miniaturized glucose **sensor** with potential for use in vivo)

RN 125275-25-4 HCAPLUS

CN 3,5,8-Trioxa-4-phosphaundec-10-en-1-aminium, 4-hydroxy-N,N,N,10-tetramethyl-9-oxo-, inner salt, 4-oxide, polymer with butyl 2-methyl-2-propenoate (9CI) (CA INDEX NAME)

CM 1

CRN 67881-98-5 CMF C11 H22 N O6 P

CM 2

CRN 97-88-1 CMF C8 H14 O2

L23 ANSWER 10 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1998:512479 HCAPLUS

DN 129:221223

ED Entered STN: 18 Aug 1998

- TI Soluble cellulose derivatives, their manufacture, grafted products, and biocompatible materials
- IN Fukui, Hiroki; Matsuyama, Kazuo; Ishihara, Kazuhiko; Nakahayashi, Nobuo
- PA Nippon Oil and Fats Co., Ltd., Japan; Nakabayashi, Norio; Foundation for Scientific Technology Promotion
- SO Jpn. Kokai Tokkyo Koho, 11 pp. CODEN: JKXXAF
- DT Patent
- LA Japanese
- IC ICM C08B015-02

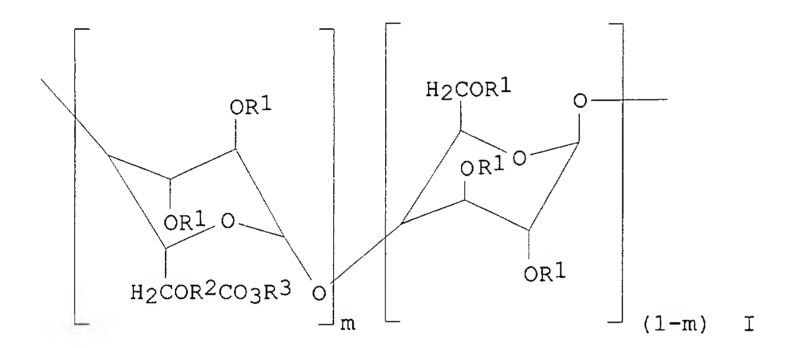
ICS A61L027-00; C08B003-22; C08F251-02

CC 63-7 (Pharmaceuticals)

Section cross-reference(s): 43

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI PRAI GI	JP 10212301 JP 1997-14988	A2	19980811 19970129	JP 1997-14988	19970129



Soluble cellulose derivs. I [R1 = H, Me, (hydroxy)ethyl, hydroxypropyl, CH2CO2H, acetyl, NO2; R2 = C1-15 hydrocarbylene; R3 = C4-15 hydrocarbyl; m = 0.001-1] are prepared by reaction of soluble celluloses with AR2CO3R3 (A = halo; R2, R3 = same as I) in the presence of bases. The biocompatible materials (e.g. hemodialyzers) contain grafted celluloses prepared by graft polymerization of radically polymerizable monomers onto I. 2- (Methacryloyloxy)ethyl 2-(trimethylammonio)ethyl phosphate was polymerized in the presence of cellulose derivative (prepared from hydroxypropyl Me cellulose and BrCH2CO3CMe3) to give a graft polymer. Cellulose membrane was coated with an aqueous solution of the graft polymer to show a low protein adsorption. ST cellulose graft polymer biocompatible medical material; hemodialyzer membrane cellulose graft polymer; peroxy ester cellulose graft polymer

IT Organ, animal

(artificial; preparation of soluble cellulose graft polymers for biocompatible

medical materials)

IT Medical goods

(catheters; preparation of soluble cellulose graft polymers for biocompatible

medical materials)

IT Dialyzers

(hemodialyzers; preparation of soluble cellulose graft polymers for

biocompatible medical materials)

IT Biosensors

(preparation of soluble cellulose graft polymers for biocompatible medical materials)

IT 868-77-9DP, 2-Hydroxyethyl methacrylate, graft copolymers with tert-butylperoxycarbonylmethyl hydroxypropyl Me cellulose 67881-98-5DP, 2-(Methacryloyloxy)ethyl 2-(trimethylammonio)ethyl phosphate, graft copolymers with tert-butylperoxycarbonylmethyl hydroxypropyl Me cellulose 87026-37-7DP, reaction products with hydroxypropyl Me cellulose, graft copolymers with 2-(methacryloyloxy)ethyl 2-(trimethylammonio)ethyl phosphate 88475-85-8DP, tert-Butylperoxy 4-(bromomethyl)benzoate, reaction products with hydroxypropyl Me cellulose, graft copolymers with 2-(methacryloyloxy)ethyl 2-(trimethylammonio)ethyl phosphate RL: PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of soluble cellulose graft polymers for biocompatible medical materials)

75-91-2, tert-Butyl hydroperoxide 9004-65-3, Hydroxypropyl methyl cellulose 22118-09-8, Bromoacetyl chloride 52780-16-2, 4-(Bromomethyl)benzoyl chloride RL: RCT (Reactant); RACT (Reactant or reagent) (preparation of soluble cellulose graft polymers for biocompatible medical

(preparation of soluble cellulose graft polymers for biocompatible medical materials)

IT 87026-37-7P 88475-85-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of soluble cellulose graft polymers for biocompatible medical materials)

Offs81-98-5DP, 2-(Methacryloyloxy)ethyl 2-(trimethylammonio)ethyl phosphate, graft copolymers with tert-butylperoxycarbonylmethyl hydroxypropyl Me cellulose
RL: PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use);
BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of soluble cellulose graft polymers for biocompatible medical materials)

RN 67881-98-5 HCAPLUS

CN 3,5,8-Trioxa-4-phosphaundec-10-en-1-aminium, 4-hydroxy-N,N,N,10-tetramethyl-9-oxo-, inner salt, 4-oxide (9CI) (CA INDEX NAME)

L23 ANSWER 11 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1996:15020 HCAPLUS

DN 124:161638

ED Entered STN: 06 Jan 1996

TI Polysiloxane-based biomembranes

AU Miller, L. S.; Rhoden, A. L.; Byrne, N.; Heptinstall, J.; Walton, D. J.

CS School Natural Environmental Sciences, Coventry University, Coventry, CV1 5FB, UK

SO Materials Science & Engineering, C: Biomimetic Materials, Sensors and Systems (1995), C3(3-4), 187-90

CODEN: MSCEEE; ISSN: 0928-4931

PB Elsevier

DT Journal

LA English

CC 76-3 (Electric Phenomena)
 Section cross-reference(s): 36

Natural phosphatidylcholine was attached to a polysiloxane backbone. The Langmuir film, Langmuir-Blodgett deposition and Montal-Mueller properties were studied. The material forms exceptionally stable Langmuir films. Montal-Mueller bilayers have a resistance of about 2 + 104 Ω cm-2 and a breakdown voltage of about 350 mV. Langmuir-Blodgett bilayer films have a low-voltage resistance of about 2 + 105 Ω cm-2, a transition to a square-law dependence at around 0.1 V, and a breakdown voltage greater than 1 V. There is evidence that bacteriorhodopsin in readily incorporated into such films.

ST polysiloxane attached phosphatidylcholine Langmuir Blodgett film; elec resistance breakdown voltage LB film; Montal Mueller bilayer polysiloxane based biomembrane

IT Dielectric strength

Electric resistance

(Langmuir film, Langmuir-Blodgett deposition and Montal-Mueller properties of polysiloxane-based biomembranes)

IT 17118-56-8DP, reaction product with dimethylsilanediolmethylsilsilanediol copolymer 156118-35-3DP,
Dimethylsilanediol-methylsilanediol copolymer, reaction product with phosphatidylcholine

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (Langmuir film, Langmuir-Blodgett deposition and Montal-Mueller properties of polysiloxane-based biomembranes)

IT 17118-56-8DP, reaction product with dimethylsilanediol-methylsilsilanediol copolymer

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (Langmuir film, Langmuir-Blodgett deposition and Montal-Mueller properties of polysiloxane-based biomembranes)

RN 17118-56-8 HCAPLUS

CN 3,5,9-Trioxa-4-phosphaheptacos-18-en-1-aminium, 4-hydroxy-N,N,N-trimethyl-10-oxo-7-[(1-oxohexadecyl)oxy]-, inner salt, 4-oxide, (18Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

Me
$$(CH_2)_{14}$$
 O $(CH_2)_{7}$ Z $(CH_2)_{7}$ Me

L23 ANSWER 12 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1994:293211 HCAPLUS

DN 120:293211

ED Entered STN: 11 Jun 1994

TI Studies on biocompatible membranes. (II). Biocompatibility of poly(acrylonitrile) copolymers with phosphonolipid polar groups

- Lee, Mi Kyung; Jung, Suk Gyu; Kim, Han Do; Cho, Hyun Hok; Kim, Kyung Hwan; AU Park, Soo Min Coll. Eng., Pusan Natl. Univ., Pusan, S. Korea CS Journal of the Korean Fiber Society (1993), 30(11), 823-8 SO CODEN: HSKCDQ; ISSN: 0253-6420 DTJournal English LA9-1 (Biochemical Methods) CC Section cross-reference(s): 13, 80 AB A methacrylate monomer having the phosphonolipid polar group,
- A methacrylate monomer having the phosphonolipid polar group, 2-(methacryloxyl)-2-(trimethylammonium) Et phosphate (MTP) was prepared and copolymd. with acrylonitrile (AN) and glycidylmethacrylate (GMA). The polymer membranes and enzyme immobilized membranes were prepared from the copolymers by a solution casting technique. Amts. of protein adsorption on polymer surface were investigated using a sorption method. Protein adsorption on the surface of the copolymer with phosphonolipid group was suppressed effectively. In platelet adhesion tests, the copolymer with a phosphonolipid group displayed less platelet adhesion than homopolymers, poly(acrylonitrile), poly(hydroxyethyl methacrylate) and Cuprophane ®. The current reproducibility of glucose sensor was obtained using various glucose solns. The current was reproducible within 40 h in the presence of plasma proteins.
- ST methacryloxyltrimethylammonium ethyl phosphate acrylonitrile glycidylmethacrylate membrane; glucose **sensor** biocompatible membrane
- IT Blood analysis

 (glucose determination in, using methacryloxyltrimethylammonium Et phosphate acrylonitrile glycidylmethacrylate biocompatible membrane sensor)
- IT Electrodes

(bio-, enzyme, glucose-selective, construction of, using methacryloxyltrimethylammonium Et phosphate acrylonitrile glycidylmethacrylate biocompatible membrane)

IT 9004-34-6, Cellulose, biological studies 25014-41-9, Polyacrylonitrile
25249-16-5, Poly(2-hydroxyethyl methacrylate)
RL: BIOL (Biological study)

(biocompatibility of)

- IT 50-99-7, Glucose, analysis
 - RL: ANT (Analyte); ANST (Analytical study)
 (determination of, using methacryloxyltrimethylammonium Et phosphate acrylonitrile glycidylmethacrylate biocompatible membrane sensor)
- IT 9001-37-0, Glucose oxidase

RL: PROC (Process)

(immobilization of, on methacryloxyltrimethylammonium Et phosphate acrylonitrile glycidylmethacrylate copolymer membrane)

- IT 154957-77-4P
 - RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and biocompatibility of, glucose sensor construction based on)
- IT 154957-77-4P
 - RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation and biocompatibility of, glucose sensor construction based on)
- RN 154957-77-4 HCAPLUS
- CN 3,5,8-Trioxa-4-phosphaundec-10-en-1-aminium, 4-hydroxy-N,N,N,10tetramethyl-9-oxo-, inner salt, 4-oxide, polymer with oxiranylmethyl 2-methyl-2-propenoate and 2-propenenitrile (9CI) (CA INDEX NAME)

PEZZUTO 10/081628 7/1/04 Page 48

CM 1

CRN 67881-98-5 CMF C11 H22 N O6 P

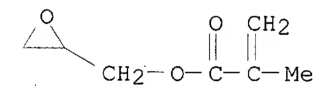
CM 2

CRN 107-13-1 CMF C3 H3 N

 $H_2C = CH - C = N$

CM 3

CRN 106-91-2 CMF C7 H10 O3



L23 ANSWER 13 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1994:26638 HCAPLUS

DN 120:26638

ED Entered STN: 22 Jan 1994

TI Polymeric liposomes formed from a new phosphatidylcholine with terminal diene groups

AU Anikin, Alexei; Chupin, Vladimir; Anikin, Michael; Serebrennikova, Galina; Tarahovsky, Jury

CS M. V. Lomonosov Inst. Fine Chem. Technol., Moscow, 117571, Russia

SO Makromolekulare Chemie (1993), 194(10), 2663-73 CODEN: MACEAK; ISSN: 0025-116X

DT Journal

LA English

CC 9-2 (Biochemical Methods)

Section cross-reference(s): 6, 23

AB A novel polymerizable phospholipid with conjugated diene groups at the hydrocarbon chain ends, 1,2-bis(11,13-tetradecadienoyl)-sn-glycero-3-phosphocholine, was prepared This phospholipid gives liposomes which can be readily polymerized upon γ - and UV-irradiation. The conversion is up to 95% within 10 h at 60° or within 3 days at room temperature. According to light scattering, 31P NMR and electron microscopy, the polymerized liposomes are detergent-resistant and maintain their structure under ultrasonic treatment and in organic solvent media.

```
polymer liposome phosphatidylcholine terminal diene prepn; membrane
ST
     phosphatidylcholine terminal diene prepn
     Kinetics of polymerization
IT
        (of bis(tetradecadienoyl)glycerophosphocholine, liposome preparation for
        diagnostic and biosensor uses in relation to)
     Membrane, biological
IT
        (polyphosphatidylcholine derivative for , preparation and properties of)
IT
     Liposome
        (multilamellar, poly[bis(tetradecadienoyl)glycerophosphocholine],
        preparation and properties of, for biosensors and diagnostics
        purposes)
     Polymerization
ΙT
        (photochem., of bis(tetradecadienoyl)glycerophosphocholine, liposome
        preparation for diagnostic and biosensor uses in relation to)
     Phosphatidylcholines, polymers
IT
     RL: PREP (Preparation)
        (polymers, preparation and properties of liposomes of, for
        biosensors and diagnostic purposes)
     Polymerization
IT
        (radiochem., of bis(tetradecadienoyl)glycerophosphocholine, liposome
        preparation for diagnostic and biosensor uses in relation to)
IT
     Polymerization
        (thermal, of bis(tetradecadienoyl)glycerophosphocholine, liposome
        preparation for diagnostic and biosensor uses in relation to)
     107-02-8, Acrolein, reactions
ΙT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (Wittig reaction of, with (methoxycarbonyldecyl)triphenylphosphonium
        bromide)
IT
     2834-05-1
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (esterification of)
IT
     68532-63-8P
     RL: RCT (Reactant); PREP (Preparation); RACT (Reactant or reagent)
        (preparation and Wittig reaction of, with acrolein)
     151416-65-8P
ΙT
     RL: RCT (Reactant); PREP (Preparation); RACT (Reactant or reagent)
        (preparation and acylation by, of glycerophosphocholinecadmium dichloride
        complex)
     151416-63-6P 151416-64-7P
     RL: PREP (Preparation)
        (preparation and conversion to anhydride)
     151416-66-9P
IT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation and homopolymn. of)
                   151416-67-0P
IT
     151416-62-5P
     RL: RCT (Reactant); PREP (Preparation); RACT (Reactant or reagent)
        (preparation and hydrolysis of)
\operatorname{IT}
     151927-35-4P
     RL: PREP (Preparation)
        (preparation and properties of liposomes of, for biosensors and
        diagnostic uses)
     6287-90-7P
IT
     RL: PREP (Preparation)
        (preparation and reaction of triphenylphosphine)
     603-35-0, Triphenylphosphine, reactions
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with Me bromoundecanoate)
     151927-35-4P
IT
     RL: PREP (Preparation)
```

(preparation and properties of liposomes of, for biosensors and diagnostic uses)

RN 151927-35-4 HCAPLUS

CN 3,5,9-Trioxa-4-phosphatricosa-20,22-dien-1-aminium, 4-hydroxy-N,N,N-trimethyl-10-oxo-7-[(1-oxo-11,13-tetradecadienyl)oxy]-, inner salt, 4-oxide, (7R)-, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 139879-59-7 CMF C36 H64 N O8 P

Absolute stereochemistry.
Double bond geometry unknown.

L23 ANSWER 14 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1992:37513 HCAPLUS

DN 116:37513

ED Entered STN: 08 Feb 1992

TI Biosensors with ion channel-containing liquid crystalline membranes

IN Gitler, Carlos; Yuli, Itzhak

PA Yeda Research and Development Co., Ltd., Israel

SO Eur. Pat. Appl., 19 pp.

CODEN: EPXXDW

DT Patent

LA English

IC ICM G01N033-543 ICS C12M001-40

CC 9-7 (Biochemical Methods)

FAN.CNT 1

FAN.CNT	1				
PAS	TENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI EP	441120	A2	19910814	EP 1991-100198	19910108
ΕP	441120	A3	19920122		
EP	441120	В1	19951129		
EP	441120	B2	20020403		
	R: AT, BE,	CH, DE	, DK, ES, FR,	GB, GR, IT, LI, LU	, NL, SE
IL	93020	A1	19950629	IL 1990-93020	19900109
CA	2033776	AA	19910710	CA 1991-2033776	19910108
CA	2033776	C	20010313		
AT	130938	E	19951215	AT 1991-100198	19910108
ES	2082867	Т3	19960401	ES 1991-100198	19910108
AU	9169245	A1	19910711	AU 1991-69245	19910109
AU	625017	В2	19920625		
US	5204239	A	19930420	US 1991-638488	19910109

```
IT
    Receptors
     RL: ANST (Analytical study)
        (cholinergic, lipid bilayer containing, biosensor containing)
IT
     Phospholipids, compounds
     RL: ANST (Analytical study)
        (conjugates, with hydrophilic spacer arm, in lipid bilayer attachment
        to biosensor)
     Phosphatidylethanolamines
IT
     RL: ANST (Analytical study)
        (conjugates, with thiol- or thioether-terminated polyoxyalkylene, for
        lipid bilayer attachment to biosensor)
IT Polyoxyalkylenes, compounds
     RL: ANST (Analytical study)
        (mercapto-terminated, conjugates, with phosphatidylethanolamines, for
        lipid bilayer attachment to biosensor)
     Antibodies
ΙT
     RL: ANST (Analytical study)
        (monoclonal, to trinitrobenzene, reactivity with trinitrobenzene-containing
        peptide conjugate with mellitin-derived peptide and lipid
        bilayer-containing membrane electrode)
     Phosphatidylethanolamines
IT
     RL: ANST (Analytical study)
        (reaction products, with dibromo PEG derivative, for lipid bilayer
        attachment to biosensor)
    138250-15-4
IT
     RL: ANST (Analytical study)
        (as ion channel, in biosensor with ion channel-containing lipid
        bilayer)
     51-84-3, Acetyl choline, analysis
IT
     RL: ANT (Analyte); ANST (Analytical study)
        (determination of, biosensor with acetylcholine receptor-containing lipid
        bilayer for)
     37231-28-0D, Melittin, peptide analogs
{f T}
    RL: ANST (Analytical study)
        (for ion channel in lipid bilayer in biosensor)
     57-88-5, Cholesterol, biological studies 59-02-9, \alpha-Tocopherol
IT
     RL: ANST (Analytical study)
        (mixed micelle containing, in ion channel-containing lipid bilayer
preparation for
       biosensor)
    76779-16-3P
ΙT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and reaction of, with phosphatidylethanolamine, for lipid
       bilayer attachment to biosensor)
     138002-99-0P
\operatorname{IT}
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, for attachment of ion channel-containing lipid bilayer to
       biosensor)
    25322-68-3
IT
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with phosphoric trichloride)
    138250-16-5
IT
    RL: PRP (Properties)
        (reactivity of, with anti-trinitrobenzene monoclonal antibody and lipid
        bilayer-containing electrode)
    138002-99-0P
IT
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, for attachment of ion channel-containing lipid bilayer to
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biosensor)

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RN 138002-99-0 HCAPLUS

CN Poly(oxy-1,2-ethanediyl), α -[2-[[2-[[(2,3-dihydroxypropoxy)hydroxyphosphinyl]oxy]ethyl]amino]ethyl]- ω -(2-mercaptoethoxy)- (9CI) (CA INDEX NAME)

OH OH OH HO-CH2-CH2-CH2-CH2-NH-CH2-CH2-CH2-O-CH2

PAGE 1-B

 $-CH_2$ $0-CH_2-CH_2-SH$

L23 ANSWER 15 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1991:234837 HCAPLUS

DN 114:234837

ED Entered STN: 15 Jun 1991

TI Vinylpyridinium copolymers as adsorbents for microorganisms

IN Yoshimatsu, Akira; Kondo, Akihiro

PA Kao Corp., Japan

SO Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

IC ICM A01N043-40

ICA C08F008-44; C08F226-06; C08L039-08

CC 61-5 (Water)

Section cross-reference(s): 10, 16, 38

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

PI JP 02237901 A2 19900920 JP 1989-59180 19890310

PRAI JP 1989-59180 19890310

Adsorbents for microorganisms contain vinylpyridinium copolymers prepared by copolymn. of vinylpyridines with hydrophilic crosslinkable monomers (and other vinyl monomers), followed by quaternization. The adsorbents have good mech. strength, quickly adsorb microorganisms from water or air or from solid materials without adversely affecting the water, air, or surface. The adsorbents are also useful for immobilization of microorganisms in bioreactors or biosensors. A suitable polymer is formed by mixing 4-vinylpyridine 105, diethylene glycol dimethacrylate 13, CaCO3 10, AIBN 1, and Me benzoate 100 g in water at 80° for 3 h to give 110 g copolymer, which (21 g) is treated with 120 g PhCH2Br in MeOH at 60° for 5 h to give 48 g 1-benzyl-4-vinylpyridinium bromide-diethylene glycol dimethacrylate copolymer (I). I at 2 g adsorbed Escherichia coli (6.5 + 107/mL dispersed in 20 mL physiol. saline solution) within 16 min, compared with 2.5 h for 1-benzyl-4-vinylpyridinium

2

bromide-divinylbenzene copolymer. microorganism adsorbent vinylpyridinium copolymer; water treatment STvinylpyridinium copolymer adsorbent; pyridinium vinyl copolymer water treatment; immobilization microorganism vinylpyridinium copolymer Air conditioning IT(vinylpyridinium copolymers as adsorbents for microorganisms in) Quaternary ammonium compounds, polymers IT RL: OCCU (Occurrence) (vinylpyridinium copolymers, as adsorbents for microorganisms in water ΙT Adsorbents (vinylpyridinium copolymers, for microorganisms removal from air or water) Wastewater treatment ITWater purification (disinfection, vinylpyridinium copolymers as adsorbents for removal of microorganisms in) IT132705-85-2P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and quaternization of) 132682-47-4P 132682-49-6P 132682-51-0P 132682-52-1P $\operatorname{\mathbf{I}}\mathbf{T}$ **132682-53-2P** 132682-54-3P 132705-86-3P 132705-87-4P 132705-89-6P 132705-91-0P 132745-98-3P 133927-76-1P RL: PREP (Preparation) (preparation of, as adsorbent for microorganisms in water or air) 100-39-0, Benzyl bromide ${ t IT}$ RL: RCT (Reactant); RACT (Reactant or reagent) (quaternization by, of vinylpyridine copolymer) 132682-53-2P ΙT RL: PREP (Preparation) (preparation of, as adsorbent for microorganisms in water or air) RN132682-53-2 HCAPLUS 2-Propenoic acid, 3,6,9,12,15,18,21-heptaoxatricosane-1,23-diyl ester, CNpolymer with 2-ethenylpyridine, compd. with (bromomethyl)benzene, phosphate (9CI) (CA INDEX NAME) CM1 CRN 7664-38-2 CMF H3 O4 P HO- b- OH OH CM2 CRN 100-39-0

CMF C7 H7 Br

CRN 132682-48-5

CMF (C22 H38 O11 . C7 H7 N) \times

CCI PMS

CM 4

CRN 134704-83-9 CMF C22 H38 O11

PAGE 1-A

PAGE 1-B

$$\begin{array}{c} \text{O} \\ \parallel \\ -\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{C}+\text{CH}_2-\text{CH}_2-\text{O}-\text{C}+\text{CH}_2-\text{CH}_2-\text{O}-\text{C}+\text{CH}_2-\text{CH}_2-\text{CH}_2-\text{O}-\text{C}+\text{CH}_2-\text{CH$$

CM 5

CRN 100-69-6 CMF C7 H7 N

L23 ANSWER 16 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1989:213278 HCAPLUS

DN 110:213278

ED Entered STN: 10 Jun 1989

TI Preparation of lipid membranes and their use

IN Ono, Seigo; Nakaya, Tadao

PA Oki Electric Industry Co., Ltd., Japan

SO Jpn. Kokai Tokkyo Koho, 6 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

IC ICM C07F009-10

ICS B32B009-00; C08F030-02

CC 33-6 (Carbohydrates)

Section cross-reference(s): 5, 7, 9, 63

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE JP 63222186 A2 19880916 PΙ JP 1987-53977 19870311 JP 06004654 B4 19940119 PRAI JP 1987-53977 19870311 CH2:CRCONHCH2CH2OP(O)(OH)OCH2CH[O2C(CH2)14Me]CH2O2C(CH2)14Me [I; R = H, Me] are polymerized to give membranes useful as ion-permeable membranes (no data), etc. H3N+CH2CH2OP(O)(O-)OCH2CH[O2C(CH2)14Me]CH2O2C(CH2)14Me was reacted with CH2:CHCOCl in pyridine containing Et3N to give I (R = H), a benzene solution of which was spread on a silicone plate, which was then irradiated with γ ray to give a white, elastic membrane. lipid membrane manuf ion permeable STAnimal tissue culture IT Biosensors Plant tissue culture (artificial lipid membranes for, preparation of) Humidity IT(sensors for, artificial lipid membranes for, preparation of) ITOrgan (artificial, artificial lipid membranes for, preparation of) 814-68-6, Acryloyl chloride 920-46-7, Methacryloyl chloride ITRL: RCT (Reactant); RACT (Reactant or reagent) (amidation by, of phosphatidylcholine) 5681-36-7 IT RL: RCT (Reactant); RACT (Reactant or reagent) (amidation of, by acryloyl chloride and methacroyl chloride) IT119845-22-6P 119845-24-8P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as ion-permeable membrane) 119845-23-7P IT119845-21-5P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, for ion-permeable membrane) IT119845-22-6P 119845-24-8P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as ion-permeable membrane) 119845-22-6 HCAPLUS RNHexadecanoic acid, 1-(3-hydroxy-3-oxido-8-oxo-2,4-dioxa-7-aza-3-phosphadec-CN 9-en-1-yl)-1,2-ethanediyl ester, homopolymer (9CI) (CA INDEX NAME) CM1 CRN 119845-21-5 CMF C40 H76 N O9 P

RN 119845-24-8 HCAPLUS

CN Hexadecanoic acid, 1-(3-hydroxy-9-methyl-3-oxido-8-oxo-2,4-dioxa-7-aza-3-phosphadec-9-en-1-yl)-1,2-ethanediyl ester, homopolymer (9CI) (CA INDEX

NAME)

CM 1

CRN 119845-23-7 CMF C41 H78 N O9 P

L23 ANSWER 17 OF 17 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1983:551151 HCAPLUS

DN 99:151151

ED Entered STN: 12 May 1984

TI Calcium ion-selective electrode studies: covalent bonding of organic phosphates and phosphonates to polymer matrixes

AU Hobby, P. C.; Moody, G. J.; Thomas, J. D. R.

CS Appl. Chem. Dep., Univ. Wales Inst. Sci. Technol., Cardiff, CF1 3NU, UK

SO Analyst (Cambridge, United Kingdom) (1983), 108(1286), 581-90 CODEN: ANALAO; ISSN: 0003-2654

DT Journal

LA English

CC 79-2 (Inorganic Analytical Chemistry) Section cross-reference(s): **35**, 72

AB The preparation and Ca2+ selectivity of organic phosphate- and phosphonate-grafted

polymer membranes were studied for partly hydrolyzed vinyl acetate-vinyl chloride polymer grafted with monodecyl phosphate (I), mono-[4-(1,1,3,3-tetramethylbutyl)phenyl] phosphate (II), and monooctyl phenylphosphonate. The preparation of phosphated polystyrene membranes by Friedel-Crafts and free-radical processes was attempted. Membranes containing I and II required phenylphosphonate solvent mediator. The electrodes had good response but poor Ca2+ selectivity compared with easily fabricated membranes of phys. mixts. of sensor plus solvent mediator in PVC.

ST calcium ion selective electrode; org phosphate polymer membrane electrode; phosphonate graft copolymer membrane; vinyl acetate chloride polymer membrane

IT Electrodes

(calcium-selective, organic phosphate- and phosphonate-grafted polymer membranes)

IT 7440-70-2, analysis

RL: ANT (Analyte); ANST (Analytical study)

(determination of, selective electrode with covalent bonding of organic phosphates

and phosphonates to polymer matrixes for)

1T 1986-91-0DP, polymers with partly hydrolyzed vinyl
acetate-vinyl chloride polymer 3921-30-0DP,
polymers with partly hydrolyzed vinyl acetate-vinyl chloride

9003-22-9DP, partly hydrolyzed, polymers with organic polymer phosphates and phosphonates 13244-67-2DP, polymers with partly hydrolyzed vinyl acetate-vinyl chloride polymer RL: PREP (Preparation) (graft, membranes, preparation and calcium ion selectivity of) 1986-91-0DP, polymers with partly hydrolyzed vinyl IT acetate-vinyl chloride polymer 3921-30-0DP, polymers with partly hydrolyzed vinyl acetate-vinyl chloride polymer 13244-67-2DP, polymers with partly hydrolyzed vinyl acetate-vinyl chloride polymer RL: PREP (Preparation) (graft, membranes, preparation and calcium ion selectivity of) 1986-91-0 HCAPLUS RNPhenol, 4-(1,1,3,3-tetramethylbutyl)-, dihydrogen phosphate (9CI) CN (CA INDEX NAME)

RN 3921-30-0 HCAPLUS CN Phosphoric acid, monodecyl ester (8CI, 9CI) (CA INDEX NAME)

 $H_{2}O_{3}PO-(CH_{2})_{9}-Me$

RN 13244-67-2 HCAPLUS CN Phosphonic acid, phenyl-, monooctyl ester (8CI, 9CI) (CA INDEX NAME)